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(71) Applicant (for all designated States except US): CREATIVE BIOMOLECULES, INC. [US/US]; 35 South Street, Hopkinton, MA 01748 (US).

(72) Inventors; and

(75) Inventors/Applicants (for US only): KUBERASAMPATH, Thangavel [IN/US]; 6 Spring Street, Medway, MA 02053 (US). OPPERMANN, Hermann [US/US]; 25 Summer Hill Road, Medway, MA 02053 (US). RUE-GER, David, C. [US/US]; 150 Edgemere Road, Apt. 4, West Roxbury, MA 02132 (US). OZKAYNAK, Engin [TR/US]; 44 Purdue Drive, Milford, MA 01757 (US).

(74) Agent: PITCHER, Edmund, R.; Lahive & Cockfield, 60 State Street, Boston, MA 02109 (US).

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(54) Title: OSTEOGENIC DEVICES

(57) Abstract

Disclosed are 1) osteogenic devices comprising a matrix containing osteogenic protein and methods of inducing endochondral bone growth in mammals using the devices; 2) amino acid sequence data, amino acid composition, solubility properties, structural features, homologies and various other data characterizing osteogenic proteins, and 3) methods of producing osteogenic proteins using recombinant DNA technology.

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OSTEOGENIC DEVICES

This invention relates to osteogenic devices, to genes encoding proteins which can induce osteogenesis in mammals and methods for their production using recombinant DNA techniques, to a method of reproducibly purifying osteogenic protein from mammalian bone, and to bone and cartilage repair procedures using the osteogenic device.

Mammalian bone tissue is known to contain one or more proteinaceous materials, presumably active during growth and natural bone healing, which can induce a developmental cascade of cellular events resulting in endochondral bone formation. This active factor (or factors) has variously been referred to in the literature as bone morphogenetic or morphogenic protein, bone inductive protein, osteogenic protein, osteogenin, or osteoinductive protein.

The developmental cascade of bone differentiation consists of recruitment of mesenchymal cells, proliferation of progenitor cells, calcification of cartilage, vascular invasion, bone formation, remodeling, and finally marrow differentiation (Reddi (1981) Collagen Rel. Res. 1:209-226).

Though the precise mechanisms underlying these phenotypic transformations are unclear, it has been shown that the natural endochondral bone differentiation activity of bone matrix can be dissociatively extracted and reconstituted with inactive residual collagenous matrix to restore full bone induction activity (Sampath and Reddi, (1981) Proc. Natl. Acad. Sci. USA 78:7599-7603). This provides an experimental method for assaying protein extracts for their ability to induce endochondral bone in vivo.

This putative bone inductive protein has been shown to have a molecular mass of less than 50 kilodaltons (kD). Several species of mammals produce closely related protein as demonstrated by cross species implant experiments (Sampath and Reddi (1983) Proc. Natl. Acad. Sci. USA 80:6591-6595).

The potential utility of these proteins has been widely recognized. It is contemplated that the availability of the protein would revolutionize orthopedic medicine, certain types of plastic surgery, and various periodontal and craniofacial reconstructive procedures.

The observed properties of these protein fractions have induced an intense research effort in various laboratories directed to isolating and identifying the pure factor or factors responsible for osteogenic activity. The current state of the art of purification of osteogenic protein from

mammalian bone is disclosed by Sampath et al. (Proc. Natl. Acad. Sci. USA (1987) 80). Urist et al. (Proc. Soc. Exp. Biol. Med. (1984) 173:194-199) disclose a human osteogenic protein fraction which was extracted from demineralized cortical bone by means of a calcium chloride-urea inorganic-organic solvent mixture, and retrieved by differential precipitation in guanidine-hydrochloride and preparative gel electrophoresis. The authors report that the protein fraction has an amino acid composition of an acidic polypeptide and a molecular weight in a range of 17-18 kD.

Urist et al. (Proc. Natl. Acad. Sci. USA (1984) 81:371-375) disclose a bovine bone morphogenetic protein extract having the properties of an acidic polypeptide and a molecular weight of approximately 18 kD. The authors reported that the protein was present in a fraction separated by hydroxyapatite chromatography, and that it induced bone formation in mouse hindquarter muscle and bone regeneration in trephine defects in rat and dog skulls. Their method of obtaining the extract from bone results in ill-defined and impure preparations.

European Patent Application Serial No. 148,155, published October 7, 1985, purports to disclose osteogenic proteins derived from bovine, porcine, and human origin. One of the proteins, designated by the inventors as a P3 protein having a molecular weight of 22-24 kD, is said to have been

purified to an essentially homogeneous state. This material is reported to induce bone formation when implanted into animals.

International Application No. PCT/087/01537, published January 14, 1988, discloses an impure fraction from bovine bone which has bone induction qualities. The named applicants also disclose putative bone inductive factors produced by recombinant DNA techniques. Four DNA sequences were retrieved from human or bovine genomic or cDNA libraries and apparently expressed in recombinant host cells. While the applicants stated that the expressed proteins may be bone morphogenic proteins, bone induction was not demonstrated. See also Urist et al., EP 0,212,474 entitled Bone Morphogenic Agents.

Wang et al. (Proc. Nat. Acad. Sci. USA (1988) 85: 9484-9488) discloses the purification of a bovine bone morphogenetic protein from guanidine extracts of demineralized bone having cartilage and bone formation activity as a basic protein corresponding to a molecular weight of 30 kD determined from gel elution. Purification of the protein yielded proteins of 30, 18 and 16 kD which, upon separation, were inactive. In view of this result, the authors acknowledged that the exact identity of the active material had not been determined.

Wozney et al. (Science (1988) 242: 1528-1534) discloses the isolation of full-length cDNA's encoding the human equivalents of three polypeptides originally purified from bovine bone. The authors report that each of the three recombinantly expressed human proteins are independently or in combination capable of inducing cartilage formation. No evidence of bone formation is reported.

It is an object of this invention to provide osteogenic devices comprising matrices containing dispersed osteogenic protein capable of bone induction in allogenic and xenogenic implants.

Another object is to provide a reproducible method of isolating osteogenic protein from mammalian bone tissue. Another object is to characterize the protein responsible for osteogenesis. Another object is to provide natural and recombinant osteogenic proteins capable of inducing endochondral bone formation in mammals, including humans. Yet another object is to provide genes encoding osteogenic proteins and methods for their production using recombinant DNA techniques. Another object is to provide methods for inducing cartilage formation.

These and other objects and features of the invention will be apparent from the description, drawings, and claims which follow.

Summary of the Invention

This invention involves osteogenic devices which, when implanted in a mammalian body, can induce at the locus of the implant the full developmental cascade of endochondral bone formation and bone marrow differentiation. Suitably modified as disclosed herein, the devices also may be used to induce cartilage formation. The devices comprise a carrier material, referred to herein as a matrix, having the characteristics disclosed below, containing dispersed osteogenic protein either in its native form as purified from natural sources or produced using recombinant DNA techniques.

Key to these developments was the successful development of a protocol which results in retrieval of active, substantially pure osteogenic protein from mammalian bone, and subsequent elucidation of amino acid sequence and structure data of native osteogenic protein. The protein has a half-maximum bone forming activity of about 0.8 to 1.0 ng per mg of implant. The protein is believed to be a dimer. It appears not to be active when reduced. Various combinations of species of the proteins may exist as heterodimers or homodimers.

The invention provides native forms of osteogenic protein, extracted from bone or produced using recombinant DNA techniques. The substantially

pure osteogenic protein may include forms having varying glycosylation patterns, varying N-termini, a family of related proteins having regions of amino acid sequence homology, and active truncated or mutated forms of native protein, no matter how derived. The naturally sourced osteogenic protein in its native form is glycosylated and has an apparent molecular weight of about 30 kD as determined by SDS-PAGE. When reduced, the 30 kD protein gives rise to two glycosylated polypeptide chains having apparent molecular weights of about 16 kD and 18 kD. In the reduced state, the 30 kD protein has no detectable osteogenic activity. The deglycosylated protein, which has osteogenic activity, has an apparent molecular weight of about 27 kD. When reduced, the 27 kD protein gives rise to the two deglycosylated polypeptides have molecular weights of about 14 kD to 16 kD.

Analysis of intact molecules and digestion fragments indicate that the native 30 kD osteogenic protein contains the following amino acid sequences (question marks indicate undetermined residues):

- (1) S-F-D-A-Y-Y-C-S-G-A-C-Q-F-P-M-P-K;
- (2) S-L-K-P-S-N-Y-A-T-I-Q-S-I-V;
- (3) A-C-C-V-P-T-E-L-S-A-I-S-M-L-Y-L-D-E-N-E-K;
- (4) M-S-S-L-S-I-L-F-F-D-E-N-K;
- (5) S-Q-E-L-Y-V-D-F-Q-R;
- (6) F-L-H-C-Q-F-S-E-R-N-S;
- (7) T-V-G-Q-L-N-E-Q-S-S-E-P-N-I-Y;

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- (8) L-Y-D-P-M-V-V;
- (9) V-G-V-V-P-G-I-P-E-P-C-C-V-P-E;
- (10) V-D-F-A-D-I-G;
- (11) V-P-K-P-C-C-A-P-T;
- (12) I-N-I-A-N-Y-L;
- (13) D-N-H-V-L-T-M-F-P-I-A-I-N;
- (14) D-E-Q-T-L-K-K-A-R-R-K-Q-W-I-?-P;
- (15) D-I-G-?-S-E-W-I-I-?-P;
- (16) S-I-V-R-A-V-G-V-P-G-I-P-E-P-?-?-V;
- (17) D-?-I-V-A-P-P-Q-Y-H-A-F-Y;
- (18) D-E-N-K-N-V-V-L-K-V-Y-P-N-M-T-V-E;
- (19) S-Q-T-L-Q-F-D-E-Q-T-L-K-?-A-R-?-K-Q;
- (20) D-E-Q-T-L-K-K-A-R-R-K-Q-W-I-E-P-R-N-?-A-R-R-Y-L;
- (21) A-R-R-K-Q-W-I-E-P-R-N-?-A-?-R-Y-?-?-V-D; and
- (22) R-?-Q-W-I-E-P-?-N-?-A-?-?-Y-L-K-V-D-?-A-?-?-G.

The availability of the protein in substantially pure form, and knowledge of its amino acid sequence and other structural features, enable the identification, cloning, and expression of native genes which encode osteogenic proteins. When properly modified after translation, incorporated in a suitable matrix, and implanted as disclosed herein, these proteins are operative to induce formation of cartilage and endochondral bone.

Consensus DNA sequences designed as disclosed herein based on partial sequence data and observed homologies with regulatory proteins disclosed in the literature are useful as probes for extracting genes encoding osteogenic protein from genomic and cDNA libraries. One of the consensus

sequences has been used to isolate a heretofore unidentified genomic DNA sequence, portions of which when ligated encode a protein having a region capable of inducing endochondral bone formation. This protein, designated OP1, has an active region having the sequence set forth below.

1 10 20 30 40
OP1 LYVSFR-DLGWQDWIIAPEGYAAYYCEGECAFPLNS
50 60 70

YMNATN--H-AIVQTLVHFINPET-VPKPCCAPTQLNA
80 90 100
ISVLYFDDSSNVILKKYRNMVVRACGCH

A longer active sequence is:

-5
HQRQA
1 10 20 30 40
OP1 CKKHELYVSFR-DLGWQDWIIAPEGYAAYYCEGECAFPLNS
50 60 70
YMNATN--H-AIVQTLVHFINPET-VPKPCCAPTQLNA
80 90 100
ISVLYFDDSSNVILKKYRNMVVRACGCH

Fig. 1A discloses the genomic DNA sequence of OP1.

The probes have also retrieved the DNA sequences identified in PCT/087/01537, referenced above, designated therein as BMPII(b) and BMPIII. The inventors herein have discovered that certain subparts of these genomic DNAs, and BMPIIa, from the same publication, when properly assembled, encode proteins (CBMPIIa, CBMPIIb, and CBMPIII) which have true osteogenic activity, i.e., induce the full cascade of events when properly implanted in a mammal

leading to endochondral bone formation. These sequences are:

CBMP-2a CKRHPLYVDFS-DVGWNDWIVAPPGYHAFYCHGECPFPLAD
50 60 70
HLNSTN--H-AIVQTLVNSVNS-K-IPKACCVPTELSA
80 90 100
ISMLYLDENEKVVLKNYQDMVVEGCGCR

1 10 20 30 40
CBMP-2b CRRHSLYVDFS-DVGWNDWIVAPPGYQAFYCHGDCPFPLAD
50 60 70
HLNSTN--H-AIVQTLVNSVNS-S-IPKACCVPTELSA
80 90 100
ISMLYLDEYDKVVLKNYQEMVVEGCGCR

1 10 20 30 40
CBMP-3 CARRYLKVDFA-DIGWSEWIISPKSFDAYYCSGACQFPMPK
50 60 70
SLKPSN--H-ATIQSIVRAVGVVPGIPEPCCVPEKMSS
80 90 100
LSILFFDENKNVVLKVYPNMTVESCACR

Thus, in view of this disclosure, skilled genetic engineers can isolate genes from cDNA or genomic libraries which encode appropriate amino acid sequences, and then can express them in various types of host cells, including both procaryotes and eucaryotes, to produce large quantities of active proteins capable of inducing bone formation in mammals including humans.

æ

The substantially pure osteogenic proteins (i.e., naturally derived or recombinant proteins free of contaminating proteins having no osteoinductive activity) are useful in clinical applications in conjunction with a suitable delivery or support system (matrix). The matrix is made up of particles or porous materials. pores must be of a dimension to permit progenitor cell migration and subsequent differentiation and proliferation. The particle size should be within the range of 70 - 850 $\mu m\text{, preferably 70 - 420 }\mu m\text{.}$ It may be fabricated by close packing particulate material into a shape spanning the bone defect, or by otherwise structuring as desired a material that is biocompatible (non-inflammatory) and, biodegradable in vivo to serve as a "temporary scaffold" and substratum for recruitment of migratory progenitor cells, and as a base for their subsequent anchoring and proliferation. Currently preferred carriers include particulate, demineralized, guanidine extracted, species-specific (allogenic) bone, and particulate, deglycosylated (or HF treated), protein extracted, demineralized, xenogenic bone. Optionally, such xenogenic bone powder matrices also may be treated with proteases such as trypsin. Other useful matrix materials comprise collagen, homopolymers and copolymers of glycolic acid and lactic acid, hydroxyapatite, tricalcium phosphate and other calcium phosphates.

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The osteogenic proteins and implantable osteogenic devices enabled and disclosed herein will permit the physician to obtain optimal predictable bone formation to correct, for example, acquired and congenital craniofacial and other skeletal or dental anomalies (Glowacki et al. (1981) Lancet 1:959-963). The devices may be used to induce local endochondral bone formation in non-union fractures as demonstrated in animal tests, and in other clinical applications including periodontal applications where bone formation is required. Another potential clinical application is in cartilage repair, for example, in the treatment of osteoarthritis.

Brief Description of the Drawing

The foregoing and other objects of this invention, the various features thereof, as well as the invention itself, may be more fully understood from the following description, when read together with the accompanying drawings, in which:

FIGURE 1A represents the nucleotide sequence of the genomic copy of osteogenic protein "OP1" gene. The unknown region between 1880 and 1920 actually represents about 1000 nucleotides;

FIGURE 1B is a representation of the hybridization of the consensus gene/probe to the osteogenic protein "OP1" gene;

FIGURE 2 is a collection of plots of protein concentration (as indicated by optical absorption) vs elution volume illustrating the results of bovine osteogenic protein (BOP) fractionation during purification on heparin-Sepharose-I; HAP-Ultragel; sieving gel (Sephacryl 300); and heparin-Sepharose-II;

FIGURE 3 is a photographic reproduction of a Coomassie blue stained SDS polyacrylamide gel of the osteogenic protein under non-reducing (A) and reducing (B) conditions;

FIGURE 4 is a photographic reproduction of a Con A blot of an SDS polyacrylamide gel showing the carbohydrate component of oxidized (A) and reduced (B) 30 kD protein;

FIGURE 5 is a photographic reproduction of an autoradiogram of an SDS polyacrylamide gel of 1251-labelled glycosylated (A) and deglycosylated (B) osteogenic protein under non-reducing (1) and reducing (2) conditions;

FIGURE 6 is a photographic reproduction of an autoradiogram of an SDS polyacrylamide gel of peptides produced upon the digestion of the 30 kD osteogenic protein with V-8 protease (B), Endo Lys C protease (C), pepsin (D), and trypsin (E). (A) is control;

FIGURE 7 is a collection of HPLC chromatograms of tryptic peptide digestions of 30 kD BOP (A), the 16 kD subunit (B), and the 18 kD subunit (C);

FIGURE 8 is an HPLC chromatogram of an elution profile on reverse phase C-18 HPLC of the samples recovered from the second heparin-Sepharose chromatography step (see FIGURE 2D). Superimposed is the percent bone formation in each fraction;

FIGURE 9 is a gel permeation chromatogram of an elution profile on TSK 3000/2000 gel of the C-18 purified osteogenic peak fraction. Superimposed is the percent bone formation in each fraction;

FIGURE 10 is a collection of graphs of protein concentration (as indicated by optical absorption) vs. elution volume illustrating the results of human protein fractionation on

heparin-Sepharose I (A), HAP-Ultragel (B), TSK 3000/2000 (C), and heparin-Sepharose II (D). Arrows indicate buffer changes;

FIGURE 11 is a graph showing representative dose response curves for bone-inducing activity in samples from various purification steps including reverse phase HPLC on C-18 (A), Heparin-Sepharose II (B), TSK 3000 (C), HAP-ultragel (D), and Heparin-Sepharose I (E);

FIGURE 12 is a bar graph of radiomorphometric analyses of feline bone defect repair after treatment with an osteogenic device (A), carrier control (B), and demineralized bone (C);

FIGURE 13 is a schematic representation of the DNA sequence and corresponding amino acid sequence of a consensus gene/probe for osteogenic protein (COPO);

FIGURE 14 is a graph of osteogenic activity vs. increasing molecular weight showing peak bone forming activity in the 30 kD region of an SDS polyacrylamide gel;

FIGURE 15 is a photographic representation of a Coomassie blue stained SDS gel showing gel purified subunits of the 30 kD protein;

FIGURE 16 is a pair of HPLC chromatograms of Endo Asp N proteinase digests of the 18 kD subunit (A) and the 16 kD subunit (B);

FIGURE 17 is a photographic representation of the histological examination of bone implants in the rat model: carrier alone (A); carrier and glycosylated osteogenic protein (B); and carrier and deglycosylated osteogenic protein (C). Arrows indicate osteoblasts;

FIGURE 18 is a graph illustrating the activity of xenogenic matrix (deglycolylated bovine matrix); and

FIGURES 19A and 19B are bar graphs showing the specific activity of naturally sourced OP before and after gel elution as measured by calcium content vs. increasing concentrations of proteins (dose curve, in ng).

Description

Purification protocols have been developed which enable isolation of the osteogenic protein present in crude protein extracts from mammalian bone. While each of the separation steps constitute a known separation technique, it has been discovered that the combination of a sequence of separations exploiting the protein's affinity for heparin and for hydroxyapatite (HAP) in the presence of a denaturant such as urea is key to isolating the pure protein from the crude extract. These critical separation steps are combined with separations on hydrophobic media, gel exclusion chromatography, and elution form SDS PAGE.

The isolation procedure enables the production of significant quantities of substantially pure osteogenic protein from any mammalian species, provided sufficient amounts of fresh bone from the species is available. The empirical development of the procedure, coupled with the availability of fresh calf bone, has enabled isolation of substantially pure bovine osteogenic protein (BOP). BOP has been characterized significantly as set forth below; its ability to induce cartilage and ultimately endochondral bone growth in cat, rabbit, and rat have been studied; it has been shown to be able to induce the full developmental cascade of bone formation previously ascribed to unknown protein or proteins in heterogeneous bone extracts; and it may be used to induce formation of endochondral bone in orthopedic defects including non-union fractures. In its native form it is a glycosylated, dimeric protein. However,

it is active in deglycosylated form. It has been partially sequenced. Its primary structure includes the amino acid sequences set forth herein.

Elucidation of the amino acid sequence of BOP enables the construction of pools of nucleic acid probes encoding peptide fragments. Also, a consensus nucleic acid sequence designed as disclosed herein based on the amino acid sequence data, inferred codons for the sequences, and observation of partial homology with known genes, also has been used as a probe. The probes may be used to isolate naturally occuring cDNAs which encode active mammalian osteogenic proteins (OP) as described below using standard hybridization methodology. The mRNAs are present in the cytoplasm of cells of various species which are known to synthesize osteogenic proteins. Useful cells harboring the mRNAs include, for example, osteoblasts from bone or osteosarcoma, hypertrophic chondrocytes, and stem cells. The mRNAs can be used to produce cDNA libraries. Alternatively, relevant DNAs encoding osteogenic protein may be retrieved from cloned genomic DNA libraries from various mammalian species.

These discoveries enable the construction of DNAs encoding totally novel, non-native protein constructs which individually, and combined are capable of producing true endochondral bone. They also permit expression of the natural material, truncated forms, muteins, analogs, fusion proteins,

and various other variants and constructs, from cDNAs retrieved from natural sources or synthesized using the techniques disclosed herein using automated, commercially available equipment. The DNAs may be expressed using well established recombinant DNA technologies in procaryotic or eucaryotic host cells, and may be oxidized and refolded in vitro if necessary for biological activity.

The isolation procedure for obtaining the protein from bone, the retrieval of an osteogenic protein gene, the design and production of recombinant protein, the nature of the matrix, and other material aspects concerning the nature, utility, how to make, and how to use the subject matter claimed herein will be further understood from the following, which constitutes the best method currently known for practicing the various aspects of the invention.

A - PURIFICATION OF BOP

Al. Preparation of Demineralized Bone

Demineralized bovine bone matrix is prepared by previously published procedures (Sampath and Reddi (1983) Proc. Natl. Acad. Sci. USA 80:6591-6595). Bovine diaphyseal bones (age 1-10 days) are obtained from a local slaughterhouse and used fresh. bones are stripped of muscle and fat, cleaned of periosteum, demarrowed by pressure with cold water, dipped in cold absolute ethanol, and stored at -20°C. They are then dried and fragmented by crushing and pulverized in a large mill. Care is taken to prevent heating by using liquid nitrogen. The pulverized bone is milled to a particle size between 70-420 µm and is defatted by two washes of approximately two hours duration with three volumes of chloroform and methanol (3:1). The particulate bone is then washed with one volume of absolute ethanol and dried over one volume of anhydrous ether. The defatted bone powder (the alternative method is to obtain Bovine Cortical Bone Powder (75-425 µm) from American Biomaterials) is then demineralized with 10 volumes of 0.5 N HCl at 4°C for 40 min., four times. Finally, neutralizing washes are done on the demineralized bone powder with a large volume of water.

A2. Dissociative Extraction and Ethanol Precipitation

Demineralized bone matrix thus prepared is dissociatively extracted with 5 volumes of 4 M guanidine-HCl (Gu-HCl), 50mM Tris-HCl, pH 7.0, containing protease inhibitors (5 mM benzamidine, 44 mM 6-aminohexanoic acid, 4.3 mM N-ethylmaleimide, 0.44 mM phenylmethylsulfonyfluoride) for 16 hr. at 4°C. The suspension is filtered. The supernatant is collected and concentrated to one volume using an ultrafiltration hollow fiber membrane (Amicon, YM-10). The concentrate is centrifuged (8,000 x g for 10 min. at 4°C), and the supernatant is then subjected to ethanol precipitation. To one volume of concentrate is added five volumes of cold (-70°C) absolute ethanol (100%), which is then kept at -70°C for 16 hrs. The precipitate is obtained upon centrifugation at 10,000 x g for 10 min. at 4°C. resulting pellet is resuspended in 4 l of 85% cold ethanol incubated for 60 min. at -70°C and recentrifuged. The precipitate is again resuspended in 85% cold ethanol (2 1), incubated at -70°C for 60 min. and centrifuged. The precipitate is then lyophilized.

A3. Heparin-Sepharose Chromatography I

The ethanol precipitated, lyophilized, extracted crude protein is dissolved in 25 volumes of 6 M urea, 50 mM Tris-HCl, pH 7.0 (Buffer A) containing 0.15 M NaCl, and clarified by centrifugation at 8,000 x g for 10 min. The

heparin-Sepharose is column-equilibrated with Buffer A. The protein is loaded onto the column and after washing with three column volume of initial buffer (Buffer A containing 0.15 M NaCl), protein is eluted with Buffer A containing 0.5 M NaCl. The absorption of the eluate is monitored continuously at 280 nm. The pool of protein eluted by 0.5 M NaCl (approximately 1 column volumes) is collected and stored at 4°C.

As shown in FIGURE 2A, most of the protein (about 95%) remains unbound. Approximately 5% of the protein is bound to the column. The unbound fraction has no bone inductive activity when bioassayed as a whole or after a partial purification through Sepharose CL-6B.

A4. Hydroxyapaptite-Ultrogel Chromatography

The volume of protein eluted by Buffer A containing 0.5 M NaCl from the heparin-Sepharose is applied directly to a column of hydroxyapaptite-ultrogel (HAP-ultrogel) (LKB Instruments), equilibrated with Buffer A containing 0.5 M NaCl. The HAP-ultrogel is treated with Buffer A containing 500 mM Na phosphate prior to equilibration. The unadsorbed protein is collected as an unbound fraction, and the column is washed with three column volumes of Buffer A containing 0.5 M NaCl. The column is subsequently eluted with Buffer A containing 100 mM Na Phosphate (FIGURE 2B).

The eluted component can induce endochondral bone as measured by alkaline phosphatase activity and histology. As the biologically active protein is bound to HAP in the presence of 6 M urea and 0.5 M NaCl, it is likely that the protein has an affinity for bone mineral and may be displaced only by phosphate ions.

A5. Sephacryl S-300 Gel Exclusion Chromatography

Sephacryl S-300 HR (High Resolution, 5 cm x 100 cm column) is obtained from Pharmacia and equilibrated with 4 M guanidine-HCl, 50 mM Tris-HCl, pH 7.0. The bound protein fraction from HA-ultrogel is concentrated and exhanged from urea to 4 M guanidine-HCl, 50 mM Tris-HCl, pH 7.0 via an Amicon ultrafiltration YM-10 membrane. The solution is then filtered with Schleicher and Schuell CENTREX disposable microfilters. A sample aliquot of approximately 15 ml containing approximately 400 mg of protein is loaded onto the column and then eluted with 4 M guanidine-HCl, 50 mM Tris-HCl, pH 7.0, with a flow rate of 3 ml/min; 12 ml fractions are collected over 8 hours and the concentration of protein is measured at A280nm (FIGURE 2C). An aliquot of the individual fractions is bioassayed for bone formation. Those fractions which have shown bone formation and migrate with an apparent molecular weight of less than 35 kD are pooled and concentrated via an Amicon ultrafiltration system with YM-10 membrane.

A6. Heparin-Sepharose Chromatography-II

The pooled osteo-inductive fractions obtained from gel exclusion chromatography are dialysed extensively against distilled water (dH₂0 and then against 6 M urea, 50 mM Tris-HCl, pH 7.0 (Buffer A) containing 0.1 M NaCl. The dialysate is then cleared through centrifugation. The sample is applied to the heparin-sepharose column (equilibrated with the same buffer). After washing with three column volumes of initial buffer, the column is developed sequentially with Buffer B containing 0.15 M NaCl, and 0.5 M NaCl (FIGURE 2D). The protein eluted by 0.5 M NaCl is collected and dialyzed extensively against distilled water. It is then dialyzed against 30% acetonitrile, 0.1% TFA at 4°C.

A7. Reverse Phase HPLC

The protein is further purified by C-18

Vydac silica-based HPLC column chromatography

(particle size 5 µm; pore size 300 A). The

osteoinductive fraction obtained from

heparin-sepharose-II chromatograph is loaded onto the

column, and washed in 0.1% TFA, 10% acetonitrile for

five min. As shown in FIGURE 8, the bound proteins

are eluted with a linear gradient of 10-30%

acetonitrile over 15 min., 30-50% acetonitrile over

60 min, and 50-70% acetonitrile over 10 min at 22°C

with a flow rate of 1.5 ml/min and 1.4 ml samples are

collected in polycarbonate tubes. Protein is

monitored by absorbance at A214 nm. Column fractions are tested for the presence of osteoinductive activity, and concanavalin A-blottable proteins. These fractions are then pooled, and characterized biochemically for the presence of 30 kD protein by autoradiography, concanavalin A blotting, and Coomassie blue dye staining. They are then assayed for in vivo osteogenic activity. Biological activity is not found in the absence of 30 kD protein.

A8. Gel Elution

The glycosylated or deglycosylated protein is eluted from SDS gels (0.5 mm thickness) for further characterization. 125I-labelled 30 kD protein is routinely added to each preparation to monitor yields. TABLE 1 shows the various elution buffers that have been tested and the yields of 125I-labelled protein.

TABLE 1
Elution of 30 kD Protein from SDS Gel

Buffer	<pre>% Eluted</pre>
(1) dH ₂ O	22
(2) 4 M Guanidine-HCl, Tris-HCl, pH 7.0	2
(3) 4 M Guanidine-HCl, Tris-HCl, pH 7.0, 0.5% Triton x 100	93
(4) 0.1% SDS, Tris-HCl, pH 7.0	98

TABLE 2 lists the steps used to isolate the 30 kD or deglycosylated 27 kD gel-bound protein. The standard protocol uses diffusion elution using 4M guanidine-HCl containing 0.5% Triton x 100 in Tris-HCl buffer or in Tris-HCl buffer containing 0.1% SDS to achieve greater than 95% elution of the protein from the 27 or 30 kD region of the gel for demonstration of osteogenic activity in vivo as described in later section.

TABLE 2 Preparation of Gel Eluted Protein

(C-18 Pool or deglycoslated protein plus 125_{I-labelled} 30 kD protein)

- 1. Dry using vacuum centrifugation;
- Wash pellet with H₂0;
- 3. Dissolve pellet in gel sample buffer (no reducing agent);
- 4. Electrophorese on pre-electrophoresed 0.5 mm mini qel;
- 5. Cut out 27 or 30 kD protein;
- Elute from gel with 0.1% SDS, 50mM Tris-HCl, pH
 7.0;
- 7. Filter through Centrex membrane;
- 8. Concentrate and wash with water in Centricon tube (10 kD membrane).

The overall yield of labelled 30 kD protein from the gel elution protocol is 50 - 60% of the loaded sample. Most of the loss occurs in the electrophoresis step, due to protein aggregation and/or smearing.

The yield is 0.5 to 1.0 μg substantially pure osteogenic protein per kg of bone.

A9. Isolation of the 16 kD and 18 kD Species

TABLE 3 summarizes the procedures involved in the preparation of the subunits. Approximately 10 μg of gel eluted 30 kD protein (FIGURE 3) is carboxymethylated and electrophoresed on an SDS-gel. The sample contains $125_{\rm I}-1{\rm abel}$ to trace yields and to use as an indicator for slicing the 16 kD and 18 kD regions from the gel. FIGURE 15 shows a Coomassie blue stained gel of gel-purified 16 kD and 18 kD proteins.

TABLE 3

Isolation of the Subunits of the 30 kD protein

(C-18 pool plus 125I-labeled 30 kD protein)

- Electrophorese on SDS gel.
- 2. Cut out 30 kD protein.
- 3. Elute with 0.1% SDS, 50 mM Tris-HCl, pH 7.0.
- 4. Concentrate and wash with H₂O in Centricon tube (10 kD membranes).
- 5. Electrophorese reduced sample on SDS gel.
- 6. Cut out the 16 kD and 18 kD subunits.
- 7. Elute with 0.1% SDS, 50 mM Tris-HCl, pH 7.0.
- 8. Concentrate and wash with H₂O in Centricon tubes.
- 9. Reduce and carboxymethylate in 1% SDS, 0.4 M Tris-HCl, pH 8.5.
- 10. Concentrate and wash with H₂O in Centricon tube.
- B. Biological Characterization of BOP

Bl. Gel Slicing:

Gel slicing experiments confirm that the isolated 30 kD protein is the protein responsible for osteogenic activity.

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Gels from the last step of the purification are sliced. Protein in each fraction is extracted in 15 mM Tris-HCl, pH 7.0 containing 0.1% SDS or in buffer containing 4 M guanidine-HCl, 0.5% non-ionic detergent (Triton x 100), 50 mM Tris-HCl. The extracted proteins are desalted, concentrated, and assayed for endochondral bone formation activity. The results are set forth in FIGURE 14. From this figure it is clear that the majority of osteogenic activity is due to protein at 30 kD region of the gel. Activity in higher molecular weight regions is apparently due to protein aggregation. These protein aggregates, when reduced, yields the 16 kD and 18 kD species discussed above.

B2. Con A-Sepharose Chromatography:

A sample containing the 30 kD protein is solubilized using 0.1% SDS, 50 mM Tris-HCl, and is applied to a column of concanavalin A (Con A)-Sepharose equilibrated with the same buffer. The bound material is eluted in SDS Tris-HCl buffer containing 0.5 M alpha-methyl mannoside. After reverse phase chromatography of both the bound and unbound fractions, Con A-bound materials, when implanted, result in extensive bone formation. Further characterization of the bound materials show a Con A-blottable 30 kD protein. Accordingly, the 30 kD glycosylated protein is responsible for the bone forming activity.

B3. Gel Permeation Chromatography:

TSK-3000/2000 gel permeation chromatography in guanidine-HCl alternately is used to achieve separation of the high specific activity fraction obtained from C-18 chromatography (FIGURE 9). The results demonstrate that the peak of bone inducing activity elutes in fractions containing substantially pure 30 kD protein by Coomassie blue staining. When this fraction is iodinated and subjected to autoradiography, a strong band at 30 kD accounts for 90% of the iodinated proteins. The fraction induces bone formation in vivo at a dose of 50 to 100 ng per implant.

B4. Structural Requirements for Biological Activity

B4-1 Activity after Digestion

Although the role of 30 kD osteogenic protein is clearly established for bone induction, through analysis of proteolytic cleavage products we have begun to search for a minimum structure that is necessary for activity in vivo. The results of cleavage experiments demonstrate that pepsin treatment fails to destroy bone inducing capacity, whereas trypsin or CNBr completely abolishes the activity.

An experiment is performed to isolate and identify pepsin digested product responsible for biological activity. The sample used for pepsin digestion was 20% - 30% pure. The buffer used is 0.1% TFA in water. The enzyme to substrate ratio is 1:10. A control sample is made without enzyme. The digestion mixture is incubated at room temperature for 16 hr. The digested product is then separated in 4 M guanidine-HCl using gel permeation chromatography, and the fractions are prepared for in vivo assay. The results demonstrate that active fractions from gel permeation chromotography of the pepsin digest correspond to peptides having an apparent molecular weight range of 8 kD - 10 kD.

B4-2 Unglycosylated Protein is Active

In order to understand the importance of the carbohydrates moiety with respect to osteogenic activity, the 30 kD protein has been chemically deglycosylated using HF (see below). After analyzing an aliquot of the reaction product by Con A blot to confirm the absence of carbohydrate, the material is assayed for its activity in vivo. The bioassay is positive (i.e., the deglycosylated protein produces a bone formation response as determined by histological examination shown in FIGURE 17C), demonstrating that exposure to HF did not destroy the biological function of the protein, and thus that the OP does not require carboyhdrate for biological activity. addition, the specific activity of the deglycosylated protein is approximately the same as that of the native glycosylated protein.

B5. Specific Activity of BOP

Experiments were performed 1) to determine the half maximal bone-inducing activity based on calcium content of the implant; 2) to estimate proteins at nanogram levels using a gel scanning method; and 3) to establish dose for half maximal bone inducing activity for gel eluted 30 kD BOP. The results demonstrate that gel eluted substantially pure 30 kD osteogenic protein induces bone at less than 5 ng per implant and exhibits half maximal bone differentiation activity at 20 ng per implant (approx. 25 mg). The purification data suggest that osteogenic protein has been purified from bovine bone to 367,307 fold after the final gel elution step with a specific activity of 47,750 bone forming units per mg of protein.

B5(a)Half Maximal Bone Differentiation Activity

The bone inducing activity is determined biochemically by the specific activity of alkaline phosphatase and calcium content of the day 12 implant. An increase in the specific activity of alkaline phosphatase indicates the onset of bone formation. Calcium content, on the other hand, is proportional to the amount of bone formed in the implant. The bone formation is therefore calculated by determining calcium content of the implant on day 12 in rats and expressed as bone forming units, which represent the amount that exhibits half maximal bone inducing activity compared to rat demineralized bone matrix. Bone induction

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exhibited by intact demineralized rat bone matrix is considered to be the maximal bone-differentiation activity for comparison.

B5(b) Protein Estimation Using Gel Scanning Techniques

A standard curve is developed employing known amounts of a standard protein, bovine serum albumin. The protein at varying concentration (50-300 ng) is loaded on a 15% SDS gel, electrophoresed, stained in comassie and destained. The gel is scanned at predetermined settings using a gel scanner at 580 nm. The area covered by the protein band is calculated and a standard curve against concentrations of protein is constructed. A sample with an unknown protein concentration is electrophoresed with BSA as a standard. The lane containing the unknown sample is scanned, and the concentration of protein is determined from the area under the curve.

B5(c)Gel Elution and Specific Activity

An aliquot of C-18 highly purified active fraction is subjected to SDS gel and sliced according to molecular weights described in FIGURE 14. Proteins are eluted from the slices in 4 M guanidine-HCl containing 0.5% Triton X-100, desalted, concentrated and assayed for endochondral bone forming activity as determined by calcium content. The C-18 highly active fractions and gel eluted

substantially pure 30 kD osteogenic protein are implanted in varying concentrations in order to determine the half maximal bone inducing activity.

FIGURE 14 shows that the bone inducing activity is due to proteins eluted in the 28-34 kD region. The recovery of activity after the gel elution step is determined by calcium content. FIGURES 19A and 19B represent the bone inducing activity for the various concentrations of 30 kD protein before and after gel elution as estimated by calcium content. The data suggest that the half maximal activity for 30 kD protein before gel elution is 69 ng per 25 mg implant and is 21 ng per 25 mg implant after elution. TABLE 4 describes the yield, total specific activity, and fold purification of osteogenic protein at each step during purification. Approximately 500 ug of heparin sepharose I fraction, 130-150 ug of the HA ultrogel fraction, 10-12 ug of the gel filtration fraction, 4-5 ug of the heparin sepharose II fraction, 0.4-0.5 ug of the C-18 highly purified fraction, and 20-25 ng of the gel eluted, substantially purified fraction is needed per 25 mg of implant for unequivocal bone formation for half maximal activity. Thus, 0.8-1.0 ng purified osteogenic protein per mg. of implant is required to exhibit half maximal bone differentiation activity in vivo.

TABLE 4
PURIFICATION OF BOP

Purification Steps	Protein (mg.)	Biological Activity Units*	Specific Activity Units/mg.	
Ethanol Precipitate**	30,000#	‡ 4, 000	0.13	1
Heparin Sepharose I	1,200#	‡ 2,400	2.00	15
HA-Ultrogel	300#	2,307	7.69	59
Gel filtration	n 20#	1,600	80.00	615
Heparin Sepharose II	5 <i>‡</i>	# 1,000	200.00	1,538
C-18 HPLC	0.070	150	2,043.00	15,715
Gel elution	0.0046	<u>a</u> 191	47,750.00	367,307

Values are calculated from 4 kg of bovine bone matrix (800 g of demineralized matrix).

^{*} One unit of bone forming activity is defined as the amount that exhibits half maximal bone differentiation activity compared to rat demineralized bone matrix, as determined by calcium content of the implant on day 12 in rats.

[#] Proteins were measured by absorbance at 280 nm.

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- @ Proteins were measured by gel scanning method compared to known standard protein, bovine serum albumin.
- ** Ethanol-precipitated guanidine extract of bovine bone is a weak inducer of bone in rats, possibly due to endogenous inhibitors. This precipitate is subjected to gel filtration and proteins less than 50 kD were separated and used for bioassay.

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C. CHEMICAL CHARACTERIZATION OF BOP

C1. Molecular Weight and Structure

Electrophoresis of the proteins after the final purification step on non-reducing SDS polyacrylamide gels reveals a diffuse band at about 30 kD as detected by both Coomassie blue staining (FIGURE 3A) and autoradiography.

In order to extend the analysis of BOP, the protein was examined under reducing conditions.

FIGURE 3B shows an SDS gel of BOP in the presence of dithiothreitol. Upon reduction, 30 kD BOP yields two species which are stained with Coomassic blue dye: a 16 kD species and an 18 kD species. Reduction causes loss of biological activity. The two reduced BOP species have been analyzed to determine if they are structurally related. Comparison of the amino acid composition and peptide mapping of the two species (as disclosed below) shows little differences, indicating that the native protein may comprise two chains having significant homology.

C2. Presence of Carbohydrate

The 30 kD protein has been tested for the presence of carbohydrate by Con A blotting after SDS-PAGE and transfer to nitrocellulose paper. The results demonstrate that the 30 kD protein has a high affinity for Con A, indicating that the protein is glycosylated (FIGURE 4A). In addition, the Con A

blots provide evidence for a substructure in the 30 kD region of the gel, suggesting heterogeneity due to varying degrees of glycosylation. After reduction (FIGURE 4B), Con A blots show evidence for two major components at 16 kD and 18 kD. In addition, it has been demonstrated that no glycosylated material remains at the 30 kD region after reduction.

In order to confirm the presence of carbohydrate and to estimate the amount of carbohydrate attached, the 30 kD protein is treated with N-glycanase, a deglycosylating enzyme with a broad specificity. Samples of the \$^{125}I\$-labelled 30 kD protein are incubated with the enzyme in the presence of SDS for 24 hours at 37°C. As observed by SDS-PAGE, the treated samples appear as a prominent species at about 27 kD (FIGURE 5A). Upon reduction, the 27 kD species is reduced to species having a molecular weight of about 14 kD - 16 kD (FIGURE 5B).

To ensure complete deglycosylation of the 30KD protein, chemical cleavage of the carbohydrate moieties using hydrogen fluoride (HF) is performed. Active osteogenic protein fractions pooled from the C-18 chromatography step are dried in vacuo over P205 in a polypropylene tube, and 50 µl freshly distilled anhydrous HF at -70°C is added. After capping the tube tightly, the mixture is kept at 0°C in an ice-bath with occasional agitation for 1 hr. The HF is then evaporated using a continuous stream of dry nitrogen gas. The tube is removed from the ice bath and the residue dried in vacuo over P205 and KOH pellets.

Following drying, the samples are dissolved in 100 µl of 50% acetonitrile/0.1% TFA and aliquoted for SDS gel analysis, Con A binding, and biological assay. Aliquots are dried and dissolved in either SDS gel sample buffer in preparation for SDS gel analysis and Con A blotting or 4 M guanidine-HCl, 50 mM Tris-HCl, pH 7.0 for biological assay.

The results show that samples are completely deglycosylated by the HF treatment: Con A blots after SDS gel electrophoreses and transfer to Immobilon membrane showed no binding of Con A to the treated samples, while untreated controls were strongly positive at 30 kD. Coomassie gels of treated samples showed the presense of a 27 kD band instead of the 30 kD band present in the untreated controls.

C3. Chemical and Enzymatic Cleavage

Cleavage reactions with CNBr are analyzed using Con A binding for detection of fragments associated with carbohydrate. Cleavage reactions are conducted using trifluoroacetic acid (TFA) in the presence and absence of CNBr. Reactions are conducted at 37°C for 18 hours, and the samples are vacuum dried. The samples are washed with water, dissolved in SDS gel sample buffer with reducing agent, boiled and applied to an SDS gel. After electrophoresis, the protein is transferred to Immobilon membrane and visualized by Con A binding. In low concentrations of acid (1%), CNBr cleaves the

majority of 16 kD and 18 kD species to one product, a species about 14 kD. In reactions using 10% TFA, a 14 kD species is observed both with and without CNBr.

Four proteolytic enzymes are used in these experiments to examine the digestion products of the 30 kD protein: 1) V-8 protease; 2) Endo Lys C protease; 3) pepsin; and 4) trypsin. Except for pepsin, the digestion buffer for the enzymes is 0.1 M ammonium bicarbonate, pH 8.3. The pepsin reactions are done in 0.1% TFA. The digestion volume is 100 ul and the ratio of enzyme to substrate is 1:10. 125I-labelled 30 kD osteogenic protein is added for detection. After incubation at 37°C for 16 hr., digestion mixtures are dried down and taken up in gel sample buffer containing dithiothreitol for SDS-PAGE. FIGURE 6 shows an autoradiograph of an SDS gel of the digestion products. The results show that under these conditions, only trypsin digests the reduced 16 kD/18 kD species completely and yields a major species at around 12 kD. Pepsin digestion yields better defined, lower molecular weight species. However, the 16 kD/18 kD fragments were not digested completely. The V-8 digest shows limited digestion with one dominant species at 16 kD.

C4. Protein Sequencing

To obtain amino acid sequence data, the protein is cleaved with trypsin or Endoproteinase Asp-N (EndoAsp-N). The tryptic digest of reduced and carboxymethylated 30 kD protein (approximately 10 µg)

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is fractionated by reverse-phase HPLC using a C-8 narrowbore column (13 cm x 2.1 mm ID) with a TFA/acetonitrile gradient and a flow rate of 150 µl/min. The gradient employs (A) 0.06% TFA in water and (B) 0.04% TFA in water and acetonitrile (1:4; v:v). The procedure was 10% B for five min., followed by a linear gradient for 70 min. to 80% B, followed by a linear gradient for 10 min. to 100% B. Fractions containing fragments as determined from the peaks in the HPLC profile (FIGURE 7A) are rechromatographed at least once under the same conditions in order to isolate single components satisfactory for sequence analysis.

The HPLC profiles of the similarly digested 16 kD and 18 kD subunits are shown in FIGURES 7B and 7C, respectively. These peptide maps are similar suggesting that the subunits are identical or are closely related.

The 16 kD and 18 kD subunits are digested with EndoAsp-N proteinase. The protein is treated with 0.5 µg EndoAsp-N in 50 mM sodium phosphate buffer, pH 7.8 at 36°C for 20 hr. The conditions for fractionation are the same as those described previously for the 30 kD, 16 kD, and 18 kD digests. The profiles obtained are shown in FIGURES 16A and 16B.

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(22)

Various peptide fragments produced using the foregoing procedures have been analyzed in an automated amino acid sequencer (Applied Biosystems 470A with 120A on-line PTH analysis). The following sequence data has been obtained:

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(I)
         S-F-D-A-Y-Y-C-S-G-A-C-Q-F-P-M-P-K;
.(2)
         S-L-K-P-S-N-Y-A-T-I-Q-S-I-V;
(3)
         A-C-C-V-P-T-E-L-S-A-I-S-M-L-Y-L-D-E-N-E-K;
 (4)
         M-S-S-L-S-I-L-F-F-D-E-N-K;
(5)
         S-Q-E-L-Y-V-D-F-Q-R;
(6)
         F-L-H-C-Q-F-S-E-R-N-S;
 (7)
         T-V-G-Q-L-N-E-Q-S-S-E-P-N-I-Y;
(8)
         L-Y-D-P-M-V-V;
(9)
         V-G-V-V-P-G-I-P-E-P-C-C-V-P-E;
(10)
         V-D-F-A-D-I-G;
         V-P-K-P-C-C-A-P-T;
(11)
(12)
         I-N-I-A-N-Y-L;
(13)
         D-N-H-V-L-T-M-F-P-I-A-I-N;
(14)
         D-E-Q-T-L-K-K-A-R-R-K-Q-W-I-?-P;
(15)
         D-I-G-?-S-E-W-I-I-?-P;
(16)
         S-I-V-R-A-V-G-V-P-G-I-P-E-P-?-?-V;
(17)
         D-?-I-V-A-P-P-Q-Y-H-A-F-Y;
(18)
         D-E-N-K-N-V-V-L-K-V-Y-P-N-M-T-V-E;
(19)
         S-Q-T-L-Q-F-D-E-Q-T-L-K-?-A-R-?-K-Q;
         D-E-Q-T-L-K-K-A-R-R-K-Q-W-I-E-P-R-N-?-A-R-R-Y
(20)
         -L;
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A-R-R-K-Q-W-I-E-P-R-N-?-A-?-R-Y-?--V-D; and R-?-Q-W-I-E-P-?-N-?-A-?--Y-L-K-V-D-?-A-?-?-G

C5. Amino Acid Analysis

Samples of oxidized (30 kD) and reduced (16 kD and 18 kD) BOP are electrophoresed on a gel and transferred to Immobilon for hydrolysis and amino acid analysis using conventional, commercially available reagents to derivatize samples and HPLC using the PicO Tag (Millipore) system. The composition data generated by amino acid analyses of 30 kD BOP is reproducible, with some variation in the number of residues for a few amino acids, especially cysteine and isoleucine.

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Composition data obtained are shown in TABLE

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TABLE 5
BOP Amino Acid Analyses

	•	•	
Amino Acid	30 kD	<u>16 kD</u>	<u>18 k</u> D
Aspartic Acid/	. 22	14	15
Asparagine			
Glutamic Acid/	24	14	16
Glutamine			
Serine	24	16	23
Glycine	29	18	26
Histidine	5	*	- 4
Arginine	13	6	6
Threonine	11	6	7
Alanine	18	11	12
Proline	14	6	6
Tyrosine	11	. 3	3
Valine	14	8	7
Methionine	3	0	2
Cysteine**	16	14	12
Isoleucine	15	14	10
Leucine	. 15	8	9
Phenylalanine	7	4	4
Tryptophan	ND	ND	ND
Lysine	12	6	6
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*This result is not integrated because histidine is present in low quantities.

**Cysteine is corrected by percent normally recovered from performic acid hydrolysis of the standard protein.

The results obtained from the 16 kD and 18 kD subunits, when combined, closely resemble the numbers obtained from the native 30 kD protein. The high figures obtained for glycine and serine are most likely the result of gel elution.

D. PURIFICATION OF HUMAN OSTEOGENIC PROTEIN

Human bone is obtained from the Bone Bank, (Massachusetts General Hospital, Boston, MA), and is milled, defatted, demarrowed and demineralized by the procedure disclosed above. 320 g of mineralized bone matrix yields 70 - 80 g of demineralized bone matrix. Dissociative extraction and ethanol precipitation of the matrix gives 12.5 g of quanidine-HCl extract.

One third of the ethanol precipitate (0.5 g) is used for gel filtration through 4 M guanidine-HCl (FIGURE 10A). Approximately 70-80 g of ethanol precipitate per run is used. In vivo bone inducing activity is localized in the fractions containing proteins in the 30 kD range. They are pooled and equilibrated in 6 M urea, 0.5 M NaCl buffer, and applied directly onto a HAP column; the bound protein

is eluted stepwise by using the same buffer containing 100 mM and 500 mM phosphate (FIGURE 10B). Bioassay of HAP bound and unbound fractions demonstrates that only the fraction eluted by 100 mM phosphate has bone inducing activity in vivo. The biologically active fraction obtained from HAP chromatography is subjected to heparin-Sepharose affinity chromatography in buffer containing low salt; the bound proteins are eluted by 0.5 M NaCl (FIGURE 10C). Assaying the heparin-Sepharose fractions shows that the bound fraction eluted by 0.5 M NaCl have bone-inducing activity. The active fraction is then subjected to C-18 reverse phase chromatography. (FIGURE 10D).

The active fraction can then be subjected to SDS-PAGE as noted above to yield a band at about 30 kD comprising substantially pure human osteogenic protein.

E. BIOSYNTHETIC PROBES FOR ISOLATION OF GENES ENCODING NATIVE OSTEOGENIC PROTEIN

E-1 PROBE DESIGN

A synthetic consensus gene shown in FIGURE 13 was designed as a hybridization probe based on amino acid predictions from homology with the TGF-beta gene family and using human codon bias as found in human TGF-beta. The designed concensus sequence was then constructed using known techniques involving assembly of oligonucleotides manufactured in a DNA synthesizer.

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Tryptic peptides derived from BOP and sequenced by Edman degradation provided amino acid sequences that showed strong homology with the Drosophila DPP protein sequence (as inferred from the gene), the Xenopus VGl protein, and somewhat less homology to inhibin and TGF-beta, as demonstrated below in TABLE 6.

TABLE 6

protein	amino acid sequence	homology
(BOP)	SFDAYYCSGACQFPS	(9/15 matches)
(DPP)	GYDAYYCHGKCPFFL	·
(<u>BOP</u>)	SFDAYYCSGACQFPS	(6/15 matches)
(<u>Vgl</u>)	GYMANYCYGECPYPL	
(BOP)	SFDAYYCSGACQFPS	(5/15 matches)
(inhibin)	GYHANYCEGECPSHI	(3) 13 maconos,
(<u>BOP</u>)	SFDAYYCSGACQFPS	(4/15 matches)
(TGF-beta)	GYHANFCLGPCPYIW	
(<u>BOP</u>)	K/RACCVPTELSAISMLYLDEN	(12/20 matches)
(<u>Vgl</u>)	LPCCVPTKMSPISMLFYDNN	

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(<u>BOP</u>) (<u>inhibin</u>)	K/RACCVPTELSAISMLYLDEN * **** * **** * KSCCVPTKLRPMSMLYYDDG	(12/20	matches)
(BOP)	K/RACCVPTELSAISMLYLDE **** * * APCCVPQALEPLPIVYYVG	(6/19	matches)
(BOP)	K/RACCVPTELSAISMLYLDEN ****** * **** KACCVPTQLDSVAMLYLNDQ	(12/20	matches)
(BOP)	LYVDF ***** LYVDF	(5/5	matches)
(BOP) (Vgl)	LYVDF *** * LYVEF	(4/5	matches)
(BOP) (TGF-beta)	LYVDF ** ** LYIDF	(4/5	matches)
(BOP)	LYVDF * * FFVSF	(2/5	matches)

^{*-}match

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In determining the amino acid sequence of an osteogenic protein (from which the nucleic acid sequence can be determined), the following points were considered: (1) the amino acid sequence determined by Edman degradation of osteogenic protein tryptic fragments is ranked highest as long as it has a strong signal and shows homology or conservative changes when aligned with the other members of the gene family; (2) where the sequence matches for all four proteins, it is used in the synthetic gene sequence; (3) matching amino acids in DPP and Vgl are used; (4) If Vgl or DPP diverged but either one were matched by inhibin or by TGF-beta, this matched amino acid is chosen; (5) where all sequences diverged, the DPP sequence is initially chosen, with a later plan of creating the Vgl sequence by mutagenesis kept as a possibility. In addition, the consensus sequence is designed to preserve the disulfide crosslinking and the apparent structural homology.

One purpose of the originally designed synthetic consensus gene sequence, designated COPO, (see FIGURE 13), was to serve as a probe to isolate natural genes. For this reason the DNA was designed using human codon bias. Alternatively, probes may be constructed using conventional techniques comprising a group of sequences of nucleotides which encode any portion of the amino acid sequence of the osteogenic protein produced in accordance with the foregoing isolation procedure. Use of such pools of probes also will enable isolation of a DNA encoding the intact protein.

E-2 Retrieval of Genes Encoding Osteogenic Protein from Genomic Library

A human genomic library (Maniatis-library) carried in lambda phage (Charon 4A) was screened using the COPO consensus gene as probe. The initial screening was of 500,000 plaques (10 plates of 50,000 each). Areas giving hybridization signal were punched out from the plates, phage particles were eluted and plated again at a density of 2000-3000 plaques per plate. A second hybridization yielded plagues which were plated once more, this time at a density of ca 100 plaques per plate allowing isolation of pure clones. The probe (COPO) is a 300 base pair BamHI-PstI fragment restricted from an amplification plasmid which was labeled using alpha 32 dCTP according to the random priming method of Feinberg and Vogelstein (1984) Anal. Biochem. 137: Prehybridization was done for 1 hr in 5x SSPE, 10x Denhardt's mix, 0.5% SDS at 50°C. Hybridization was overnight in the same solution as above plus probe. The washing of nitrocellulose membranes was done, once cold for 5 min. in 1x SSPE with 0.1% SDS and twice at 50°C for 2 x 30 min. in the same solution. Using this procedure, twenty-four positive clones were found. Two contained a gene never before reported designated OP1, osteogenic protein-I described below. Two others yielded the genes corresponding to BMP-2b, one yielded BMP-3 (see PCT US 87/01537).

Southern blot analysis of lambda #13 DNA showed that an approximately 3kb BamHI fragment hybridized to the probe. (See FIGURE 1B). fragment was isolated and subcloned into a bluescript vector (at the BamHI site). The clone was further analyzed by Southern blotting and hybridization to the COPO probe. This showed that a 1 kb (approx.) EcoRI fragment strongly hybridized to the probe. This fragment was subcloned into the EcoRI site of a bluescript vector, and sequenced. Analysis of this sequence showed that the fragment encoded the carboxy terminus of a protein, named osteogenic protein-1 (OP1). The protein was identified by amino acid homology with the TGF-beta family. For this comparison cysteine patterns were used and then the adjacent amino acids were compared. Consensus splice signals were found where amino acid homologies ended, designating exon intron boundaries. Three exons were combined to obtain a functional TGF-beta-like domain containing seven cysteines. Two introns were deleted by looping out via primers bridging the exons using the single stranded mutagenesis method of Kunkel. Also, upstream of the first cysteine, an EcoRI site and an asp-pro junction for acid cleavage were introduced, and at the 3' end a PstI site was added by the same technique. Further sequence information (penultimate exon) was obtained by sequencing the entire insert. The sequencing was done by generating a set of unidirectionally deleted clones (Ozkaynak, E., and Putney, S. (1987) Biotechniques, 5:770-773). The obtained sequence covers about 80% of the TGF-beta-like region of OP1 and is set forth in FIGURE 1A. The complete sequence of the TGF-beta like region was obtained by first subcloning all EcoRI generated fragments of lambda clone #13 DNA and sequencing a 4 kb fragment that includes the first portion of the TGF-beta like region (third exon counting from end) as well as sequences characterized earlier. The gene on an EcoRI to PstI fragment was inserted into an E. coli expression vector controlled by the trp promoter-operator to produce a modified trp LE fusion protein with an acid cleavage site. The OPl gene encodes amino acids corresponding substantially to a peptide found in sequences of naturally sourced material. The amino acid sequence of what is believed to be its active region is set forth below:

1 10 20 30 40
OP1 LYVSFR-DLGWQDWIIAPEGYAAYYCEGECAFPLNS
50 60 70

YMNATN--H-AIVQTLVHFINPET-VPKPCCAPTQLNA
80 90 100
ISVLYFDDSSNVILKKYRNMVVRACGCH

A longer active sequence is:

-5
HQRQA
1 10 20 30 40
OP1 CKKHELYVSFR-DLGWQDWIIAPEGYAAYYCEGECAFPLNS
50 60 70
YMNATN--H-AIVQTLVHFINPET-VPKPCCAPTQLNA
80 90 100
ISVLYFDDSSNVILKKYRNMVVRACGCH

The amino acid sequence of what is believed to be the active regions encoded by the other three native genes retrieved using the consensus probe are:

- CBMP-2a CKRHPLYVDFS-DVGWNDWIVAPPGYHAFYCHGECPFPLAD

 50 60 70

 HLNSTN--H-AIVQTLVNSVNS-K-IPKACCVPTELSA

 80 90 100

 ISMLYLDENEKVVLKNYQDMVVEGCGCR
- CBMP-2b 1 10 20 30 40
 CRRHSLYVDFS-DVGWNDWIVAPPGYQAFYCHGDCPFPLAD
 50 60 70
 HLNSTN--H-AIVQTLVNSVNS-S-IPKACCVPTELSA
 80 90 100
 ISMLYLDEYDKVVLKNYQEMVVEGCGCR
- CBMP-3

 1 10 20 30 40

 CARRYLKVDFA-DIGWSEWIISPKSFDAYYCSGACQFPMPK

 50 60 70

 SLKPSN--H-ATIQSIVRAVGVVPGIPEPCCVPEKMSS

 80 90 100

 LSILFFDENKNVVLKVYPNMTVESCACR

E-3 Probing cDNA Library

Another example of the use of pools of probes to enable isolation of a DNA encoding the

intact protein is shown by the following. Cells known to express the protein (e.g., osteoblasts or osteosarcoma) are extracted to isolate total cytoplasmic RNA. An oligo-dT column can be used to isolate mRNA. This mRNA can be size fractionated by, for example, gel electrophoresis. The fraction which includes the mRNA of interest may be determined by inducing transient expression in a suitable host cell and testing for the presence of osteogenic protein using, for example, antibody raised against peptides derived from the tryptic fragments of osteogenic protein in an immunoassay. The mRNA fraction is then reverse transcribed to single stranded cDNA using reverse transcriptase; a second complementary DNA strand can then be synthesized using the cDNA as a template. The double-standard DNA is then ligated into vectors which are used to transfect bacteria to produce a cDNA library.

The radiolabelled consensus sequence, portions thereof, and/or synthetic deoxy oligonucleotides complementary to codons for the known amino acid sequences in the osteogenic protein may be used to identify which of the DNAs in the cDNA library encode the full length osteogenic protein by standard DNA-DNA hybridization techniques.

The cDNA may then be integrated in an expression vector and transfected into an appropriate host cell for protein expression. The host may be a prokaryotic or eucaryotic cell since the former's inability to glycosylate osteogenic protein will not effect the protein's enzymatic activity. Useful host cells include Saccharomyces, E. coli, and various mammalian cell cultures. The vector may additionally encode various signal sequences for protein secretion and/or may encode osteogenic protein as a fusion protein. After being translated, protein may be purified from the cells or recovered from the culture medium.

E4. Gene Preparation

Natural gene sequences and cDNAs retrieved as described above may be used for expression. The genes above may also be produced by assembly of chemically synthesized oligonucleotides. 15-100mer oligonucleotides may be synthesized on a Biosearch DNA Model 8600 Synthesizer, and purified by polyacrylamide gel electrophoresis (PAGE) in Tris-Borate-EDTA buffer (TBE). The DNA is then electroeluted from the gel. Overlapping oligomers may be phosphorylated by T4 polynucleotide kinase and ligated into larger blocks which may also be purified by PAGE.

E5. Expression

The genes can be expressed in appropriate prokaryotic hosts such as various strains of E. coli, and also in bacillus, yeasts, and various animal cells such as CHO, myeloma, etc. Generally, expression may be achieved using many cell types and expression systems well known to those skilled in the art. For example, if the gene is to be expressed in E. coli, an expression vector based on pBR322 and containing a synthetic trp promoter operator and the modified trp LE leader may be used. The vector can be opened at the EcoRI and PSTI restriction sites, and, for example, an OP gene fragment can be inserted between these two sites. The OP protein is joined to the leader protein via a hinge region having the sequence Asp-Pro. This hinge permits chemical cleavage of the fusion protein with dilute acid at the Asp-Pro site.

E6. Production of Active Proteins

The following procedure may be followed for production of active recombinant proteins. E. coli cells containing the fusion proteins are lysed. The fusion proteins are purified by differential solubilization. Cleavage is conducted with dilute acid, and the resulting cleavage products are passed through a Sephacryl-200HR or SP Trisacyl column to separate the cleaved proteins. The reduced OP fractions are then subjected to HPLC on a semi-prep C-18 column.

Conditions for refolding of OP were at pH 8.0 using 50 mM Tris-HCl and 6M Gu-HCl. Samples were refolded for 18 hours at 4°C.

These procedures have been used to express in <u>E. coli</u> on the active protein designated OPl having the amino acid sequence set forth above (longer species).

Refolding may not be required if the proteins are expressed in animal cells.

MATRIX PREPARATION

A. General Consideration of Matrix Properties

The carrier described in the bioassay section, infra, may be replaced by either a biodegradable-synthetic or synthetic-inorganic matrix (e.g., HAP, collagen, tricalcium phosphate, or polylactic acid, polyglycolic acid and various copolymers thereof). Also xenogeneic bone may be used if pretreated as described below.

Studies have shown that surface charge, particle size, the presence of mineral, and the methodology for combining matrix and osteogenic protein all play a role in achieving successful bone induction. Perturbation of the charge by chemical

modification abolishes the inductive response. Particle size influences the quantitative response of new bone; particles between 75 and 420 µm elicit the maximum response. Contamination of the matrix with bone mineral will inhibit bone formation. Most importantly, the procedures used to formulate osteogenic protein onto the matrix are extremely sensitive to the physical and chemical state of both the osteogenic protein and the matrix.

The sequential cellular reactions at the interface of the bone matrix/OP implants are complex. The multistep cascade includes: binding of fibrin and fibronectin to implanted matrix, chemotaxis of cells, proliferation of fibroblasts, differentiation into chondroblasts, cartilage formation, vascular invasion, bone formation, remodeling, and bone marrow differentiation.

Must perform several important functions. It must bind osteogenic protein and act as a slow release delivery system, accommodate each step of the cellular response during bone development, and protect the osteogenic protein from nonspecific proteolysis. In addition, selected materials must be biocompatible in vivo and biodegradable; the carrier must act as a temporary scaffold until replaced completely by new bone. Biocompatibility requires that the matrix not induce significant inflammation when implanted and not be rejected by the host

animal. Biodegradability requires that the matrix be slowly absorbed by the body of the host during development of new bone or cartilage. Polylactic acid (PLA), polyglycolic acid (PGA), and various combinations have different dissolution rates in vivo. In bones, the dissolution rates can vary according to whether the implant is placed in cortical or trabecular bone.

Matrix geometry, particle size, the presence of surface charge, and porosity or the presence of interstices among the particles of a size sufficient to permit cell infiltration, are all important to successful matrix performance. It is preferred to shape the matrix to the desired form of the new bone and to have dimensions which span non-union defects. Rat studies show that the new bone is formed essentially having the dimensions of the device implanted.

The matrix may comprise a shape-retaining solid made of loosely adhered particulate material, e.g., with collagen. It may also comprise a molded, porous solid, or simply an aggregation of close-packed particles held in place by surrounding tissue. Masticated muscle or other tissue may also be used. Large allogeneic bone implants can act as a carrier for the matrix if their marrow cavities are cleaned and packed with particles and the dispersed osteogenic protein.

B. <u>Preparation of Biologically Active Allogenic</u> Matrix

Demineralized bone matrix is prepared from the dehydrated diaphyseal shafts of rat femur and tibia as described herein to produce a bone particle size which pass through a 420 μ sieve. particles are subjected to dissociative extraction with 4 M guanidine-HCl. Such treatment results in a complete loss of the inherent ability of the bone matrix to induce endochondral bone differentiation. The remaining insoluble material is used to fabricate the matrix. The material is mostly collagenous in nature, and upon implantation, does not induce cartilage and bone. All new preparations are tested for mineral content and false positives before use. The total loss of biological activity of bone matrix is restored when an active osteoinductive protein fraction or a pure protein is reconstituted with the biologically inactive insoluble collagenous matrix. The osteoinductive protein can be obtained from any vertebrate, e.g., bovine, porcine, monkey, or human, or produced using recombinant DNA techniques.

C. <u>Preparation of Deglycosylated Bone Matrix</u> for Use in Xenogenic Implant

When osteogenic protein is reconstituted with collagenous bone matrix from other species and implanted in rat, no bone is formed. This suggests that while the osteogenic protein is xenogenic (not

species specific), the matrix is species specific and cannot be implanted cross species perhaps due to intrinsic immunogenic or inhibitory components. Thus, heretofore, for bone-based matrices, in order for the osteogenic protein to exhibit its full bone inducing activity, a species specific collagenous bone matrix was required.

The major component of all bone matrices is Type I collagen. In addition to collagen, extracted bone includes non-collagenous proteins which may account for 5% of its mass. Many non-collagenous components of bone matrix are glycoproteins. Although the biological significance of the glycoproteins in bone formation is not known, they may present themselves as potent antigens by virtue of their carbohydrate content and may constitute immunogenic and/or inhibitory components that are present in xenogenic matrix.

It has now been discovered that a collagenous bone matrix may be used as a carrier to effect bone inducing activity in xenogenic implants, if one first removes the immonogenic and inhibitory components from the matrix. The matrix is deglycosglated chemically using, for example, hydrogen fluoride to achieve this purpose.

Bovine bone residue prepared as described above is sieved, and particles of the $74-420~\mu\mathrm{M}$ are collected. The sample is dried in vacuo over P_2O_5 , transferred to the reaction vessel and anhydrous hydrogen fluoride (HF) (10-20 ml/g of matrix) is then distilled onto the sample at -70°C. The vessel is allowed to warm to 0°C and the reaction mixture is stirred at this temperature for 120 min. After evaporation of the HF in vacuo, the residue is dried thoroughly in vacuo over KOH pellets to remove any remaining traces of acid.

Extent of deglycosylation can be determined from carbohydrate analysis of matrix samples taken before and after treatment with HF, after washing the samples appropriately to remove non-covalently bound carbohydrates.

The deglycosylated bone matrix is next treated as set forth below:

- 1) suspend in TBS (Tris-buffered Saline)
 1g/200 ml and stir at 4°C for 2 hrs or
 in 6 M urea, 50 mM Tris-HCl, 500 mM
 NaCl, pH 7.0 (UTBS), and stir at RT for
 30 min.;
- 2) centrifuge and wash with TBS or UTBS as in step 1); and
- 3) centrifuge; discard supernatant; water wash residue; and then lyophilize.

FABRICATION OF OSTEOGENIC DEVICE

Fabrication of osteogenic devices using any of the matrices set forth above with any of the osteogenic proteins described above may be performed as follows.

A. Ethanol precipitation

In this procedure, matrix was added to osteogenic protein in guanidine-HCl. Samples were vortexed and incubated at a low temperature. Samples were then further vortexed. Cold absolute ethanol was added to the mixture which was then stirred and incubated. After centrifugation (microfuge high speed) the supernatant was discarded. The reconstituted matrix was washed with cold concentrated ethanol in water and then lyophilized.

B. <u>Acetonitrile Trifluoroacetic Acid</u> <u>Lyophilization</u>

In this procedure, osteogenic protein in an acetonitrile trifluroacetic acid (ACN/TFA) solution was added to the carrier. Samples were vigorously vortexed many times and then lyophilized.

C. <u>Urea Lyophilization</u>

For those proteins that are prepared in urea buffer, the protein is mixed with the matrix, vortexed many times, and then lyophilized. The lyophilized material may be used "as is" for implants.

IN VIVO RAT BIOASSAY

Substantially pure BOP, BOP-rich extracts comprising protein having the properties set forth above, and several of the recombinant proteins have been incorporated in matrices to produce osteogenic devices, and assayed in rat for endochondral bone. Studies in rats show the osteogenic effect to be dependent on the dose of osteogenic protein dispersed in the osteogenic device. No activity is observed if the matrix is implanted alone. The following sets forth guidelines for how the osteogenic devices disclosed herein might be assayed for determining active fractions of osteogenic protein when employing the isolation procedure of the invention, and evaluating protein constructs and matrices for biological activity.

A. Subcutaneous Implantation

The bioassay for bone induction as described by Sampath and Reddi (Proc. Natl. Acad. Sci. USA (1983) 80: 6591-6595), herein incorporated by

reference, is used to monitor the purification protocols for endochondral bone differentiation activity. This assay consists of implanting the test samples in subcutaneous sites in allogeneic recipient rats under ether anesthesia. Male Long-Evans rats, aged 28-32 days, were used. A vertical incision (1 cm) is made under sterile conditions in the skin over the thoraic region, and a pocket is prepared by blunt dissection. Approximately 25 mg of the test sample is implanted deep into the pocket and the incision is closed with a metallic skin clip. The day of implantation is designated as day of the experiment. Implants were removed on day 12. The heterotropic site allows for the study of bone induction without the possible ambiguities resulting from the use of orthotopic sites.

B. <u>Cellular Events</u>

The implant model in rats exhibits a controlled progression through the stages of matrix induced endochondral bone development including: (1) transient infiltration by polymorphonuclear leukocytes on day one; (2) mesenchymal cell migration and proliferation on days two and three; (3) chondrocyte appearance on days five and six; (4) cartilage matrix formation on day seven; (5) cartilage calcification on day eight; (6) vascular invasion, appearance of osteoblasts, and formation of new bone on days nine and ten; (7) appearance of osteoblastic and bone remodeling and dissolution of

the implanted matrix on days twelve to eighteen; and (8) hematopoietic bone marrow differentiation in the ossicle on day twenty-one. The results show that the shape of the new bone conforms to the shape of the implanted matrix.

C. <u>Histological Evaluation</u>

Histological sectioning and staining is preferred to determine the extent of osteogenesis in the implants. Implants are fixed in Bouins Solution, embedded in parafilm, cut into 6-8 mm sections. Staining with toluidine blue or hemotoxylin/eosin demonstrates clearly the ultimate development of endochondrial bone. Twelve day implants are usually sufficient to determine whether the implants show bone inducing activity.

D. <u>Biological Markers</u>

Alkaline phosphatase activity may be used as a marker for osteogenesis. The enzyme activity may be determined spectrophotometrically after homogenization of the implant. The activity peaks at 9-10 days in vivo and thereafter slowly declines. Implants showing no bone development by histology should have little or no alkaline phosphatase activity under these assay conditions. The assay is useful for quantitation and obtaining an estimate of bone formation very quickly after the implants are removed from the rat. Alternatively the amount of bone formation can be determined by measuring the calcium content of the implant.

Implants containing osteogenic protein at several levels of purity have been tested to determine the most effective dose/purity level, in order to seek a formulation which could be produced on a commercial scale. The results are measured by specific acivity of alkaline phosphatase, calcium content, and histological examination. As noted previously, the specific activity of alkaline phosphatase is elevated during onset of bone formation and then declines. On the other hand, calcium content is directly proportional to the total amount of bone that is formed. The osteogenic activity due to osteogenic protein is represented by "bone forming units". For example, one bone forming unit represents the amount of protein that is needed for half maximal bone forming activity as compared to rat demineralized bone matrix as control and determined by calcium content of the implant on day 12.

E. Results

Dose curves are constructed for bone inducing activity in vivo at each step of the purification scheme by assaying various concentrations of protein. FIGURE 11 shows representative dose curves in rats as determined by alkaline phosphatase. Similar results are obtained when represented as bone forming units.

Approximately 10-12 µg of the TSK-fraction, 3-4

μg of heparin-Sepharose-II fraction, 0.4-0.5 μg of the C-18 column purified fraction, and 20-25 ng of gel eluted highly purified 30 kD protein is needed for unequivocal bone formation (half maximum activity). 20-25 ng of the substantially pure protein per 25 mg of implant is normally sufficient to produce endochondral bone. Thus, 1-2 ng osteogenic protein per mg of implant is a reasonable dosage, although higher dosages may be used. (See section IB5 on specific activity of osteogenic protein.)

OP1 expressed as set forth above (longer version), when assayed for activity histologically, induced cartilage and bone formation as evidenced by the presence of numerous chondrocytes in many areas of the implant and by the presence of osteoblasts surrounding vascular endothelium forming new matrix.

Deglycosylated xenogenic collagenous bone matrix (example: bovine) has been used instead of allogenic collagenous matrix to prepare osteogenic devices (see previous section) and bioassayed in rat for bone inducing activity in vivo. The results demonstrate that xenogenic collagenous bone matrix after chemical deglycosylation induces successful endochondral bone formation (FIGURE 19). As shown by specific activity of alkaline phosphotase, it is evident that the deglycosylated xenogenic matrix induced bone whereas untreated bovine matrix did not.

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Histological evaluation of implants suggests that the deglycosylated bovine matrix not only has induced bone in a way comparable to the rat residue matrix but also has advanced the developmental stages that are involved in endochondral bone differentiation. Compared to rat residue as control, the HF treated bovine matrix contains extensively remodeled bone. Ossicles are formed that are already filled with bone marrow elements by 12 days. This profound action as elicited by deglycosylated bovine matrix in supporting bone induction is reproducible and is dose dependent with varying concentration of osteogenic protein.

ANIMAL EFFICACY STUDIES

Substantially pure osteogenic protein from bovine bone (BOP), BOP-rich osteogenic fractions having the properties set forth above, and several recombinant proteins have been incorporated in matrices to produce osteogenic devices. The efficacy of bone-inducing potential of these devices was tested in cat and rabbit models, and found to be potent inducers of osteogenesis, ultimately resulting in formation of mineralized bone. The following sets forth guidelines as to how the osteogenic devices disclosed herein might be used in a clinical setting.

A. Feline Model

The purpose of this study is to establish a large animal efficacy model for the testing of the

osteogenic devices of the invention, and to characterize repair of massive bone defects and simulated fracture non-union encountered frequently in the practice of orthopedic surgery. The study is designed to evaluate whether implants of osteogenic protein with a carrier can enhance the regeneration of bone following injury and major reconstructive surgery by use of this large mammal model. The first step in this study design consists of the surgical preparation of a femoral osteotomy defect which, without further intervention, would consistently progress to non-union of the simulated fracture defect. The effects of implants of osteogenic devices into the created bone defects were evaluated by the following study protocol.

A-1. Procedure

Sixteen adult cats weighing less than 10 lbs. undergo unilateral preparation of a 1 cm bone defect in the right femur through a lateral surgical approach. In other experiments, a 2 cm bone defect was created. The femur is immediately internally fixed by lateral placement of an 8-hole plate to preserve the exact dimensions of the defect. There are three different types of materials implanted in the surgically created feline femoral defects: group I (n = 3) is a control group which undergo the same plate fixation with implants of 4 M guanidine-HCl-treated (inactivated) feline demineralized bone matrix powder (Gu-HCl-DBM) (360 mg); group II (n = 3) is a positive control group

implanted with biologically active feline demineralized bone matrix powder (DBM) (360 mg); and group III (n = 10) undergo a procedure identical to groups I-II, with the addition of osteogenic protein onto each of the Gu-HCl-DBM carrier samples. To summarize, the group III osteogenic protein-treated animals are implanted with exactly the same material as the group I animals, but with the singular addition of osteogenic protein.

All animals are allowed to ambulate ad libitum within their cages post-operatively. All cats are injected with tetracycline (25 mg/kg SQ each week for four weeks) for bone labelling. All but four group III animals are sacrificed four months after femoral osteotomy.

A-2. Radiomorphometrics

In vivo radiomorphometric studies are carried out immediately post-op at 4, 8, 12 and 16 weeks by taking a standardized x-ray of the lightly anesthesized animal positioned in a cushioned x-ray jig designed to consistently produce a true anterio-posterior view of the femur and the osteotomy site. All x-rays are taken in exactly the same fashion and in exactly the same position on each animal. Bone repair is calculated as a function of mineralization by means of random point analysis. A final specimen radiographic study of the excised bone is taken in two planes after sacrifice. X-ray

results are shown in FIGURE 12, and displaced as percent of bone defect repair. To summarize, at 16 weeks, 60% of the group III femors are united with average 86% bone defect regeneration. By contrast, the group I Gu-HCl-DMB negative-control implants exhibit no bone growth at four weeks, less than 10% at eight and 12 weeks, and 16% (± 10%) at 16 weeks with one of the five exhibiting a small amount of bridging bone. The group II DMB positive-control implants exhibited 18% (± 3%) repair at four weeks, 35% at eight weeks, 50% (± 10%) at twelve weeks and 70% (± 12%) by 16 weeks, a statistical difference of p <0.01 compared to osteogenic protein at every month. One of the three (33%) is united at 16 weeks.

A-3. Biomechanics

Excised test and normal femurs are immediately studied by bone densitometry, wrapped in two layers of saline-soaked towels, placed in two sealed plastic bags, and stored at -20° C until further study. Bone repair strength, load to failure, and work to failure are tested by loading to failure on a specially designed steel 4-point bending jig attached to an Instron testing machine to quantitate bone strength, stiffness, energy absorbed and deformation to failure. The study of test femurs and normal femurs yield the bone strength (load) in pounds and work to failure in joules. Normal femurs exhibit a strength of 96 (± 12) pounds. osteogenic protein-implanted femurs exhibited 35 (± 4) pounds, but when corrected for surface area at the site of

fracture (due to the "hourglass" shape of the bone defect repair) this correlated closely with normal bone strength. Only one demineralized bone specimen was available for testing with a strength of 25 pounds, but, again, the strength correlated closely with normal bone when corrected for fracture surface area.

A-4. Histomorphometry/Histology

Following biomechanical testing the bones are immediately sliced into two longitudinal sections at the defect site, weighed, and the volume measured. One-half is fixed for standard calcified bone histomorphometrics with fluorescent stain incorporation evaluation, and one-half is fixed for decalcified hemotoxylin/eosin stain histology preparation.

A-5. Biochemistry

Selected specimens from the bone repair site (n=6) are homogenized in cold 0.15 M NaCl, 3 mM NaHCO3, pH 9.0 by a Spex freezer mill. The alkaline phosphatase activity of the supernatant and total calcium content of the acid soluble fraction of sediment are then determined.

A-6. Histopathology

The final autopsy reports reveal no unusual or pathologic findings noted at necropsy of any of the animals studied. Portion of all major organs are

preserved for further study. A histopathological evaluation is performed on samples of the following organs: heart, lung, liver, both kidneys, spleen, both adrenals, lymph nodes, left and right quadriceps muscles at mid-femur (adjacent to defect site in experimental femur). No unusual or pathological lesions are seen in any of the tissues. Mild lesions seen in the quadriceps muscles are compatible with healing responses to the surgical manipulation at the defect site. Pulmonary edema is attributable to the euthanasia procedure. There is no evidence of any general systemic effects or any effects on the specific organs examined.

A-7. Feline Study Summary

The 1 cm and 2 cm femoral defect cat studies demonstrate that devices comprising a matrix containing disposed osteogenic protein can: (1) repair a weight-bearing bone defect in a large animal; (2) consistently induces bone formation shortly following (less than two weeks) implantation; and (3) induce bone by endochondral ossification, with a strength equal to normal bone, on a volume for volume basis. Furthermore, all animals remained healthy during the study and showed no evidence of clinical or histological laboratory reaction to the implanted device. In this bone defect model, there was little or no healing at control bone implant sites. The results provide evidence for the successful use of osteogenic devices to repair large, non-union bone defects.

B. Rabbit Model:

Bl. Procedure and Results

The purpose of this study is to establish a model in which there is minimal or no bone growth in the control animals, so that when bone induction is tested, only a strongly inductive substance will yield a positive result. Defects of 1.5 cm are created in the ulnae of rabbits with implantation of osteogenic devices or no implant.

Eight mature (greater than 10 lbs) New Zealand White rabbits with epiphyseal closure documented by X-ray were studied. Of these eight animals (one animal each was sacrificed at one and two weeks), 11 ulnae defects are followed for the full course of the eight week study. In all cases (n = 7) following osteoperiosteal bone resection, the no implant animals establish no radiographic union by eight weeks. All no implant animals develop a thin "shell" of bone growing from surrounding bone present at four weeks and, to a slightly greater degree, by eight weeks. In all cases (n = 4), radiographic union with marked bone induction is established in the osteogenic protein-implanted animals by eight weeks. As opposed to the no implant repairs, this bone repair is in the site of the removed bone.

Radiomorphometric analysis reveal 90% osteogenic protein-implant bone repair and 18% no-implant bone repair at sacrifice at eight weeks. At autopsy, the osteogenic protein bone appears normal, while "no implant" bone sites have only a soft fibrous tissue with no evidence of cartilage or bone repair in the defect site.

B-2. Allograft Device

In another experiment, the marrow cavity of the 1.5 cm ulnar defect is packed with activated osteogenic protein rabbit bone powder and the bones are allografted in an intercalary fashion. The two control ulnae are not healed by eight weeks and reveal the classic "ivory" appearance. In distinct contrast, the osteogenic protein-treated implants "disappear" radiographically by four weeks with the start of remineralization by six to eight weeks. These allografts heal at each end with mild proliferative bone formation by eight weeks.

This type of device serves to accelerate allograph repair.

B-3. Summary

These studies of 1.5 cm osteo-periosteal defects in the ulnae of mature rabbits show that: (1) it is a suitable model for the study of bone growth; (2) "no implant" or Gu-HCl negative control implants yield a small amount of periosteal-type bone, but not

medullary or cortical bone growth; (3) osteogenic protein-implanted rabbits exhibited proliferative bone growth in a fashion highly different from the control groups; (4) initial studies show that the bones exhibit 50% of normal bone strength (100% of normal correlated vol:vol) at only eight weeks after creation of the surgical defect; and (5) osteogenic protein-allograft studies reveal a marked effect upon both the allograft and bone healing.

The invention may be embodied in other specific forms without departing from the spirit or essential characteristics thereof. The present embodiments are therefore to be considered in all respects as illustrative and not restrictive, the scope of the invention being indicated by the appended claims rather than by the foregoing description, and all changes which come within the meaning and range of equivalency of the claims are therefore intended to be embraced therein.

What is claimed is:

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<u>Claims</u>

1. An osteogenic device for implantation in a mammal, said device comprising:

a biocompatible, in vivo biodegradable matrix defining pores of a dimension sufficient to permit influx, proliferation and differentiation of migratory progenitor cells from the body of said mammal; and

substantially pure osteogenic protein capable of inducing endochondral bone formation in said mammal disposed in said matrix and accessible to said cells.

- 2. Substantially pure osteogenic protein capable of inducing endochondral bone formation in a mammal when disposed within a matrix implanted in said mammal.
- 3. The device of claim 1 wherein said matrix comprises close-packed particulate matter having a particle size within the range of 70-420 $\mu m\,.$
- 4. The device of claim 1 wherein said matrix comprises demineralized, protein-extracted, particulate, allogenic bone.
- 5. The device of claim 1 wherein said matrix comprises collagen, hydroxyapatite, tricalcium phosphate, polymers comprising lactic acid monomer units, polymers comprising glycolic acid monomer units, demineralized, guanidine-extracted allogenic bone, or a mixture thereof.

- 6. The device of claim 1 wherein said matrix is shaped to span a non-union fracture in said mammal.
- 7. The device of claim 1 disposed within the marrow cavity of allogenic bone.
- 8. The device of claim 1 wherein said matrix comprises demineralized, protein extracted, particulate, deglycosylated xenogenic bone.
- 9. The device of claim 8 wherein said matrix is treated with a protease.
- 10. The invention of claim 1 or 2 wherein said osteogenic protein is unglycosylated.
- 11. The invention of claim 10 wherein said osteogenic protein has an apparent molecular weight of about 27 kD when oxidized as determined by comparison to molecular weight standards in SDS-polyacrylamide gel electrophoresis.
- 12. The invention of claim 1 or 2 wherein said osteogenic protein is glycosylated.
- 13. The invention of claim 12 wherein said osteogenic protein has an apparent molecular weight of about 30 kD when oxidized as determined by comparison to molecular weight standards in SDS-polyacrylamide gel electrophoresis.

- 14. The invention of claim 1 or 2 wherein said osteogenic protein comprises a pair of polypeptide chains.
- 15. The invention of claim 14 wherein one chain of said pair of polypeptide chains has an apparent molecular weight of about 14 kD and the other has an apparent molecular weight of about 16 kD, both as determined after reduction by comparison to molecular weight standards in SDS-polyacrylamide gel electrophoresis.
- 16. The invention of claim 14 wherein one chain of said pair of polypeptide chains has an apparent molecular weight of about 16 kD and the other has an apparent molecular weight of about 18 kD, both as determined after reduction by comparison to molecular weight standards in SDS-polyacrylamide gel electrophoresis.
- 17. The invention of claim 1 or 2 wherein said osteogenic protein has the approximate amino acid composition set forth below:

Amino acid <u>residue</u>	Rel. no. res./molec.	Amino acid <u>residue</u>	Rel. no. res./molec.
Aspartic acid/	22	Tyrosine	11
Asparagine		Valine	14
Glutamic acid/	24	Methionine	3
Glutamine		Cysteine	16
Serine	24	Isoleucine	15

Glycine	29	Leucine	15
Histidine	5	Proline	14
Arginine	13	Phenylalanine	7
Threonine	11	Tryptophan	ND
Alanine	18		
Lysine	12		

18. The invention of claim 1 or 2 wherein said osteogenic protein comprises the amino acid sequence:

VPKPCCAPT.

- 19. The invention of claim 1 or 2 wherein the half maximum bone inducing activity of said protein is 0.8 to 1.0 ng per mg of said matrix.
- 20. A method of inducing local cartilage or bone formation in a mammal comprising the step of implanting the device of claim 1 in said mammal at a locus accessible to migratory progenitor cells of said mammal.
- 21. A method of inducing endochondral bone formation in a mammal comprising the step of implanting the device of claim 1 in said mammal at a locus accessible to migratory progenitor cells of said mammal.
- 22. A method of inducing endochondral bone formation in a non-union fracture in a mammal comprising the step of implanting in the fracture in said mammal the device of claim 6.

23. The invention of claim 1 or 2 wherein the protein comprises the sequence:

OPI LYVSFR-DLGWQDWIIAPEGYAAYYCEGECAFPLNS
50 60 70
YMNATN--H-AIVQTLVHFINPET-VPKPCCAPTQLNA
80 90 100
ISVLYFDDSSNVILKKYRNMVVRACGCH

24. The device of claim 1 or 2 wherein the protein comprises the sequence:

-5
HQRQA

1 10 20 30 40

OP1 CKKHELYVSFR-DLGWQDWIIAPEGYAAYYCEGECAFPLNS
50 60 70
YMNATN--H-AIVQTLVHFINPET-VPKPCCAPTQLNA
80 90 100
ISVLYFDDSSNVILKKYRNMVVRACGCH

25. The device of claim 1 or 2 wherein the protein comprises the sequence:

1 10 20 30 40
CBMP-2a CKRHPLYVDFS-DVGWNDWIVAPPGYHAFYCHGECPFPLAD
50 60 70
HLNSTN--H-AIVQTLVNSVNS-K-IPKACCVPTELSA
80 90 100
ISMLYLDENEKVVLKNYQDMVVEGCGCR

- 26. The device of claim 1 or 2 wherein the protein comprises the sequence:
- 1 10 20 30 40
 CBMP-2b CRRHSLYVDFS-DVGWNDWIVAPPGYQAFYCHGDCPFPLAD
 50 60 70
 HLNSTN--H-AIVQTLVNSVNS-S-IPKACCVPTELSA
 80 90 100
 ISMLYLDEYDKVVLKNYQEMVVEGCGCR
- 27. The device of claim 1 or 2 wherein the protein comprises the sequence:
- CBMP-3

 CARRYLKVDFA-DIGWSEWIISPKSFDAYYCSGACQFPMPK

 50
 60
 70

 SLKPSN--H-ATIQSIVRAVGVVPGIPEPCCVPEKMSS
 80
 90
 100

 LSILFFDENKNVVLKVYPNMTVESCACR
- 28. A DNA sequence encoding an amino acid sequence sufficiently duplicative of that of the sequence encoded by the gene of FIGURE 1A such that said encoded sequence induces bone or cartilage formation when implanted in a mammal in association with a matrix.
- 29. The DNA of claim 28 encoding the same amino acid sequence as the gene set forth in FIGURE 1A.

-5

30.	The D	NA seg	uence	of	claim	28	encoding:	•
	1	10			20		30	40
OP1		LYVSE	R-DLG	WQDV	WIIAPE(SYA	AYYCEGECAF:	PLNS
			50	_	60		70	
	YMNA	TNH-	AIVQT	LVH	FINPET-	-VP	KPCCAPTQLN.	A
-		80	9	•		00		
•	ISVI	YFDDSS	NVILK	KŸRI	NMVVRA	CGC	H	

31. The DNA sequence of claim 28 encoding:

HQRQA

1 10 20 30 40

CKKHELYVSFR-DLGWQDWIIAPEGYAAYYCEGECAFPLNS

50 60 70

YMNATN--H-AIVQTLVHFINPET-VPKPCCAPTQLNA

80 90 100

ISVLYFDDSSNVILKKYRNMVVRACGCH

- 32. A cell line engineered to express the protein of claim 2.
- 33. The device of claim 1 wherein said matrix comprises demineralized, protein extracted, particulate xenogenic bone treated with HF.

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ന്	1A-1 ₁₀	20	30	40	50	. 09	70
	GGAGGTATAGGAGCTCTCTTCGATTTTAGCAAACCAGGAGTCCGAAGATCTAAGGAGAGCTGGGGGTTTGACTCC Bg l i i	CTTCGATTTT	AGCAAACCAGG	AGTCCGAAGATC: Bglii	TCTAAGGAGA II	\GCTGGGGGTT'	TGACTCC
	85	95	105	115	125	135	145
	GAGAGCTCGAGCAGTCCCCAAGACCTGGTCTTGACTCACGAGTTAGACTCCACTCAGAGGCTGACTGTCTCCAGG	CCAAGACCTG	STCTTGACTCA	CGAGTTAGAC	TCCACTCAGA	GGCTGACTGT	CTCCAGG
	SacI	PflMI					
	XhoI	TthlllI					
	160	170	180	190	200	210	220
	GICTACACCTCTAAGGGCGACACTGGGCTCAAGCAGACTGCCGTTTTCTATATGGGATGAGCCTTCACAGGGCAG	CGACACTGGG	CTCAAGCAGAC	TGCCGTTTTC	TATATGGGA	GAGCCTTCAC	AGGGCAG
	235	245	255	265	275	285	295
	CCAGTTGGGATGGGTTGAGGTTTGGCTGTAGACATCAGAAACCCAAGTCAAATGCGCTTCAACCAGTAGAAATT	AGGTTTGGCT	STAGACATCAG	AAACCCAAGI	CANATGCGC	LTCAACCAGTA	GAAAATT
	310	320	330	340	350	360	370
	CACCAGCCGCAGAGCTAAGGTTGGGTGGACATTAGGGTTGGTT	AAGGTTGGGT	3GACATTAGGG	TTGGTTGATC	CAGGAGCTCA	AACAGTGTCCT	CTGAGCC
					SacI		
	385	395	405	415	425	435	445
	CCAGCTCCTTCTGCCCCACCCACCATCTTCAGTGCTGCTTCCTCTAAGGCCACAGCTGTAGTTGGCCAGGGGG	ACCCCACCAT	CTTCAGTGCTG	CTTCCTCTCA	AGGCCACAG(CTGTAGTTGGC	CAGGGGG
					PvuII	II Ball	
							BglI
	460	470	480	490	200	510	520
	GCTTCATTATTTTTGCTCCTGGGCAGTAGGAGGAAGAATGAAT	TCCTGGGCAG	TAGGAGGAAGA	GAATGAATG	CTCTCCATG	SGTCTTTCTTA	GGAATGT
					NCOI		
	535 545 555 565 575 585 595	545	555	565	575	585	595
	GGGAACTTTTCCAGAA	GTCTCTATGT	CTTTTAGTTTG	TGTTGGGTC1	ACTTGCCCTT(CCTGAACCACT	TCCTGAC
	610	620	630	640	650	099	670
	TCCTGGACAGGATGTGCACTGAGCTTAGCTTTGGGGATCTAATAGTGACTTTACAAAGCCTCTTTGAGAAGG	ACTGATGAGCTT	LTAGCTTTGGG o I	GATCTAATAC	STGACTTTACA	AAAGCCTCTTT	GAGAAGG
	685	695	705	715	725	735	745
	TGACATTGGAACCAAGGCTTGAGCAGACACAACAAGATTGCAGGGGGGCATTGCAGGTGGAGGAAACGGCAC	CTTGAGCAGA	CACAACAAAGA	TTGCAGGGA	SGGCATTGC	AGGTGGAGGAA	ACGGCAC
					BS	BspMI-	
	760	770	780	790	800	810	820
	ATGCAAGAGCCCTGCGTGGGAGTGAGCTTGGTGTTTGGTCAATCAGTTGTCAGAGCACACCGGGCCCTGTCAGCA	GGGAGTGAGC	rtggtgtttg	STCAATCAGT	rgtcagagca	CACCGGGCCCT	GTCAGCA

ſή	1A-2 ₈₃₅	845	855	865	875	885	895
	GGCACAGCCTGGCCTGCTCTG	CTCTGAGTATC	SACAGAGAGCC	CCTGGGAAGI	TGTAGGTGGA	AGTATGACAGAGCCCCTGGGAAGTTGTAGGTGGAGGAAAGACAGGTCATGA	STCATGA
	910	920	930	940	950	096	970
	CTAGGAAAAAAGCAATCCCTCTGTTGTGGGGTGGAAGGATGCTTGCAGTGTGTGT	CCTCTGTTGTC	SGGGTGGAAGG	AAGGTTGCAG	TGTGTGTGAG	AGAGAGACAA	SACAGAC
	985	995	1005	1015	1025	1035	1045
	AGACAGACACTTCTCAATGTTT	IGTTTACAAG	IGCTCAGGCCC	TGACCCGAAT	GCTTCCAAAT	ACAAGTGCTCAGGCCCTGACCCGAATGCTTCCAAATTTACGTAGTTCTGGAAA	CTGGAAA
			Eco0	BsmI+	+	8 I	
	1060	1070	1080	1090	1100	1110	1120
	ACCCCCTGTATCATTTTCACTACTCCAAAGAAACCTCGGGAGTGTTTTTTTT	CACTACTCAAA	AGAAACCTCGG	GAGTGTTTTC	TTCTGAAAG	TCATCAGGTT	TTGACTC
	1135	1145	1155	1165	1175	1185	1195
	TCTGCTGTCTCTTCTTCTTGCTGGTGGTGGTGGTTGCTTGTCCCAGGCCCTGTCCCGCATCCTCTTGCCCCCCAGGCCCTGTCCTTTGCTTGC	rcttgctggt	GTGGTGATG	TECTTGEC	CAGGCCCTG1 EcoO	CCCGCATCCT	CITIGOCO
	1210	1220	1230	1240	1250	1260	1270
	CTGCAGAGGGATGAGTGTGTGGGGCCTCACGAGTTGAGGTTGTTCATAAGCAGATCTCTTTGAGCAGGCGCCCT	TGTTGGGGCC	CACGAGTTG	GGTTGTTCAT	PAAGCAGATC	CTTTGAGCAG	GGCGCCT
	PstI	EC00			Bglii		NarI Ps
	1285	1295	1305	1315	1325	1335	1345
	GCAGTGGCCTTGTGTGAGGCTGGAGGGGTTTCGATTCCCTTATGGAATCCAGGCAGATGTAGCATTTAAACAACA	GGCTGGAGGG	STTTCGATTCC	CTTATGGAAT	rccaggcaga ₁	IGTAGCATTTA	AACAACA
	tI .					DraI	
	1360	1370	1380	1390	1400	1410	1420
	CACGTGTATAAAAGAAACCAG	CCAGTGTCCG	CAGAAGGTTCC	CAGAAAGTATT	CATGGGATAA	CATGAG	AGAGGAA
	1435	1445	1455	1465	1475	1485	1495
	TGGGGCATTGGCACCTCCCTTAGTAGGGCCTTTGCTGGGGGTAGAAATGAGTTTTAAGGCAGGTTAGACCCTCGA	CCTTAGTAGG	SCCITIGCIG	SGGGTAGAAA	rgagttttaa(SGCAGGTTAGA	CCCTCGA
		EcoO					
	1510	1520	1530	1540	1550	1560	1570
	ACTGGCTTTTGAATCGGGAAA1	GAAATTTACC	CCCAGCCGT	CTGTGCTTC	ATTGCTGTTCA	FITACCCCCCAGCCGTTCTGTGCTTCATTGCTGTTCACATCACTGCCTAAGATG	TAAGATG
	1585	1595	1605	1615	1625	1635	1645
	GAGGAACTITGATGTGTGTGTTTTCTTTTCTCCTCACTGGGCTCTGCTTCTTCACTTCCTTGTCAATGCAGAAA	GTGTGTTTCT	<i>LTCTCCTCAC!</i>	regerrage	LTCTTCACTT	CTTGTCAATG	CAGAGAA
	1660	1670	1680	1690	1700	1710	1720
	CAGCAGCAGCACCAGAGGCAGGCCTTGTAAGAAGCACGAGCTGTATGTCAGCTTCCGAGACCTGGGCTGGCAGG	GGCAGGCCTT	STAAGAAGCA	GAGCTGTAT	STCAGCTTCC	BAGACCTGGGC	TGGCAGG
		StuI					BspMI
	1735	1745	1755	1765	1775	1785	1795
	TAAGGGGCTGGCTGGGTCTGTCTTGGGTGTGGGCCCTCTGGCGTGGGCTCCCACAGGCAGCGGGTGCTGTGCTCA	CIGICITGGG	rereeeccer	creecereee	CTCCCACAGG	CAGCGGGTGCT	GTGCTCA
			ApaI				
			ECCO				

AATHURE CUEET

1870 CTAAGGA 1945 ATTATGA 2020 GTGACCA Dr	2095 ACACAGC N	2170 AAATGGG	2245 TCCCACT	2320 TGAGAGG	2395 TCGTGAG	2470 CTCTCTT 2545 TTCCCTC	2620 GGTGTGG BS 2695 TCAGCCC
TACTTGGC TGATGCCZ	15 CCAGAAGA	0 TTTTTAGE	2235	GGGACCAT	15 ACCTTGG1	TCCCACAC	.0 2 ZATCTTGGC 15 2 GTAGATG1
1850 1860 1870 CTCGCTAGGGAAGGGTACTTGGCTAAGGA 1925 1935 1945 GGGAGCCAGCATGGGTGATGCCATTATGA 2000 2010 2020 GTTTAAGGTGAACTGCCAGTGTGACCA	2085 SCGTTCTGTC	2160 FTTTATGTTT	2235 AGCTTTGACG	0 2310 CCATTCTTTTG	2385 SAGGCTTTCA	2460 IGCATGGTTT 2535 IGAGGGGGAG	2610 TGTCACGCCA 2685 GGGGCCTGAG
1850 3CCTCGCTA 1925 GGGAGCC 2000 FTGTTTAAC	2075 CCTGTGCCC	2150 CATTTTA	2225 CTCACCGCA	2300 CTTCTTTCC	2375 CATGGTCT	2450 CAGGCTTCT 2525 FACTACTGT	2600 CTGGTGGGI
1840 1915 1990 ATGGAGGT	2065 GAGCTCCTTC Saci	2140 TTGTCCTT	2215 GATCATAGO	2290 GAGTGGTC	2365 TCAGAAGG	2440 GCTGTGCT(2515 ACGCGCGC	2590 2590 TGCAGACG(2665 CTGCTGCC
1810	CCTAGTGGGGTAGAGCTGATGCCTCACACCGGAGCTCCTTCCT	CATGGATGTCCATTTTAGGATCAGCCAAGCCCCGTCTTGTCCTTCATTTTTATGTTTTTTAGAAATGGG	GTCTTGCTCTGTCACCCAGGCTGGGTGCAGTGGTGTGATCATAGCTCACCGCAGCTTTGACGCCGTCTTCCCACT T+b111T	0 2280 2320 2310 2320 TAGGCCAAGACTATAGAGTGGTCCTTCTTTCCATTCTTTTGGGACCATGAGAGG	2335 2345 2355 2365 2375 2385 2395 CCACCCATGTTTCCTGCCCTGCTCAGAAGGCATGGTCTGAGGCTTTCACTTGGTCGTGAG	2410 2420 2430 2440 2450 2450 2460 2470 CCTTCGTGGTGGTTTCTTTCAGCATGGGGTTGGGATGCTGTGGCTTCTGCATGGTTTCCCACACTCTTT 2485 2505 2515 2525 2535 2545 CTCCTCCTCAGGACTGGATCGCGCCTGAAGGCTACGCGCGCTACTTGTGAGGGGGAGTGTGCCTTCCCTC	TGAACTCCTACATGAACGCCACCACCGCCATCGTGCAGCCTGGTGGGTG
1820 GCCAGTTAA(1895 ROX. 1000 1970 AGGTGGGCA	2045 ATGATTGCC	2120 GATCAGCCA	2195 GGCTGGGTG	2270 CTATAGGCC	2345 CCTGCTGGGC ApaI	2420 TCAGCATGG 2495 TCATCGCGC	2570 CCACCAACCA 2645 CTGCGGGGC
CATCTCTGCC CATCTCTGCC CATCTCTGCC CAPPROX CATCTGGCAGC CATCTGGCAGC	2 AGAGCTGA	.0 ATTTTAGG	15 2 TCACCCAG	260 2 AAGCTTGGAC	TCCTGCCC	2 GTTTCTTT 15 2 GACTGGAT	10 2 SATGAACGC SATGAACGC
1810 1820 GTCTTGTTTCTCATCTCTGCCA 1885 1895 TACAGGG(APPROX. 1960 1970 GTTATTAGCCTCTCTGGCAGGT Bg1I BSpMI-	2035 FAGTGGGGTA	2110 FGGATGTCCA	2185 CTTGCTCTGT	2260 227 CAGTCTACTAAGCTTGGACTA	2335 ACCCATGTTT	2410 242 CCTTCGTGGTGTTTCTTTCA 2485 249 CTCCTCCTCAGGACTGGATCA	TGAACTCCTACATGAACGCCA 2635 TCACCTGGGCCGGGCAGGCTG EII BglI
GTCTTC TACAGC GTTATT	CCTA CCTA alli MI	CAT	GTC	CAC	CC	CC.	TGAA TCAC

FIG. 1A-4 2710 2720 2730 2740 2750 2760 2770 ATTGCCATGTCATGACTTTTGGGGCCCCTTGCGCCGTTAAAAAAAA	2720 FTTTGGGGGCC ApaI ECOO 2795	2730 CCTTGCGCCG1 2805	2740 FTAAAAAAAAA 2815	2750 TCAAAAATTG 2825	2760 STACTTTATGA 2835	2770 CTGGTTT 2845
Drai 2860 2870 2880 2890 2900 2910 2920 ATGGGCTCCTTTGAGGATGCTTATTGTGGGTGCTGGTTACGGTGCCTAAGAGCACTGGGCCCCTGCTTCA Apai	2870 ATGCTTGTAGT	2880 ATTGTGGGTG	2890 CTGGTTACGG	DraI 2900 FGCCTAAGAGC	2910 CACTGGGCCCC Apai	2920 TGCT
2935 2935 2945 2955 2965 2975 2985 2995	2945 ACAGGTAAACA	2955 GATGAGAAAT	2965 rtcagtgagg	2975 3GCACAGTGA1	2985 rcagaagegege	2995 CCAG
3010	3020	3030	3040	3050	3060	3070
GATAATGGGATGGAGAGATGA	SATGAGTGGGG	GGGACCCATGGGC Ecoo NcoI	CATTTCAAGT	raaatt t cagi	GTGGGGACCCATGGGCCATTTCAAGTTAAATTTCAGTCGGGTCACCAGGAAGAT Ecoo NcoI	GGAAG2
3085	3095	3105	3115	3125	3135	3145
TCCATGTGATAATGAGATTAA	ATTAACGTGCC 3170	CAGTCACGGC	GACACTCAGT	AGGTGTTATTC 3200	CGTGCCCAGTCACGCGACACTCAGTAGGTGTTATTCCTGCTCTGCCAACAGCA 3180 3180 3220	AACAGC 3220
ACCATAGTTGATAAGAGCTGT	SCTGTTAGGGA	TTTTGTCCTT	TTGCTTAGAA	CCAAGGITCA	TAGGGATTTTGTCCTTTGCTTAGAATCCAAGGTTCAAGGACCTTGGTTATGTA	TTATGI

GCTCCCTGTCATGAACATCATCTGAGCCTTTCCTGCCTACTGATCATCCACCCTGCCTTGAATGCTTCTAGTGAC

BsmI+

Eco0

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FIG.	FIG. 1A-53610	3610	3620	3630	3640	3650	3660	3670
	TGCCAGT	TGCCAGTATCCCCTGGAAGGTGCCAGGCATGTCTCCCCGGCTGCCAGGGGACACATCTCTATCCTTCTCCAACCC	GGTGCCAGGC	ATGICICCCC	GGCTGCCAGG	GGACACATCT	CTATCCTTCT	CCAACCC
		3685	3695	3705	3715	3725	3735	3745
	CTGCCTT	CTGCCTTCATGGCCCATGGAACAGGAGTGCCATCGCCTGTGTGCACCTACTTCCATCAGTATTTCACCAGAGAT	GAACAGGAGT	GCCATCGCCC	TGTGTGCACC	TACTTCCATC	AGTATTCAC	CAGAGAT
	BglI	NCOI			ApaLI			BglI
	ı	3760	3770	3780	3790	3800	3810	3820
	CTGCAGG	CTGCAGGATCAAAGTGAATTCTCCAGGGATTGTGAAATGATGCGATTGTGGTCATGTTTAAAAAGGGGGCAACTGT	NTTCTCCAGGG	ATTGTGAAAT	GATGCGATTC	TGGTCATGTT	TAAAAGGGGG	CAACTGT
	Ι	ECORI	RI			Dr	DraI	
	PstI						•	
		3835	3845	3855	3865	3875	3885	3895
	CTTCTAG	CTTCTAGAGAGTCCTGATGAAATGCTTCCAGAGGAAATGAGCTGATGGCTGGAATTTGCTTTAAAATCATTCAAG	FGAAATGCTTC	CAGAGGAAAT	GAGCTGATGC	CTGGAATTTG	CTTTAAAATC	ATTCAAG
	XbaI						DraI	
		3910	3920	3930	3940	3950	3960	3970
	GTGGAGC	GTGGAGCAGGTGGGGAAGGGTATGGATGTGTAAGAGTTTGAAATTGTCCATCATAAAATGTGTAAAAAĠCATGCT	SGTATGGATG	TCTAAGAGTT	TGAAATTGT	CATCATAAAA	TGTGTAAAAA	ŚCATGCT
	•							

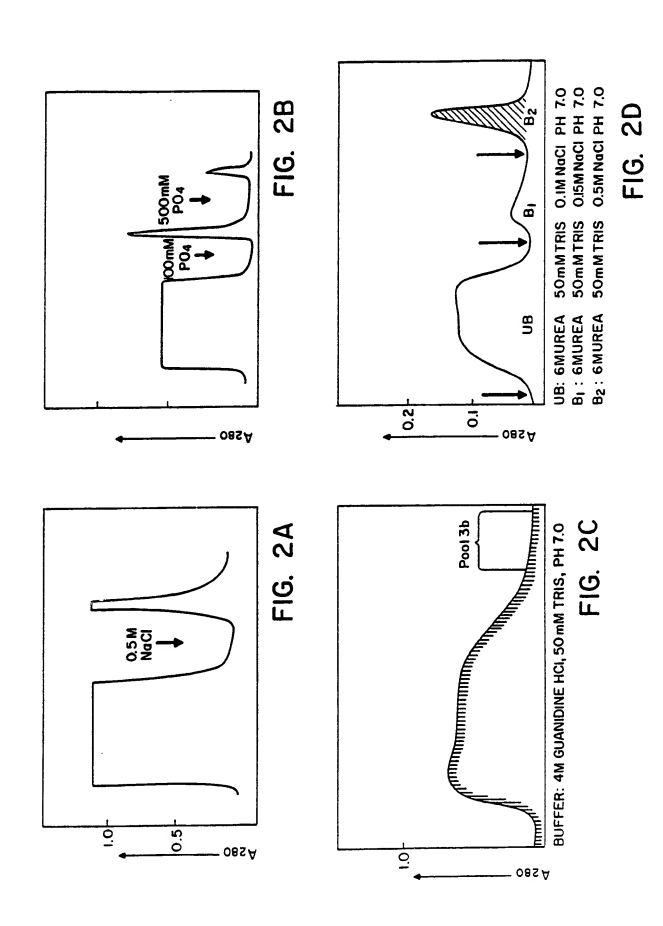
GTGGAGCAGGAAGGTATGGATTTGAAATTGTCCATCATAAAAGCATGCT BSPNI BSSNI BSNI B																						
GTGGAGCAGGTGGGAAGGTATGGATGTGAAATTGTCCATCATAAAATGTGTAA BSPMI BSPMI 3985 3985 4005 4015 4025 4035 GGCCTATGTCAGCAGTCGTAACAGTTCACATCATAAAATGTGTAA 4060 4070 4080 4080 4100 4110 4145 4145 4145 4145 4145 414	3310	AAAAĞCATGCT	SphI	4045	SGCACCTACAG	4120	SAACACGTGGC	4195	STTTGCTTAGT		4270	CATAGGCATTT	4345	CACTICTAGIG	4420	ACCTGGTCCTG		4495	LTGTCTGCAAA	4570	GGAGTGTCAC	BS
STATO STAT	3200	AAAATGTGTAA		4035	GCTCAAGCCT	4110	GCCCTGTCTT	4185	TCTTCCCTTA		4260	ACGAGGCACT	4335	CAAGATGGAG	4410	ACAGAGAATT		4485	TTTGTTCCCC	4560	GCTTGGCTTG	
GTGGAGCAGGGAATGGATGTGTAAGAGTTTGAAAJ BSPMI— 3985 3985 4005 4015 GGCCTATGTCAGCAGTCACAGGTGTGAACAGTGCC 4060 4070 4080 4090 TTGCTGGAAACCCAGAGTTTCACGTTGAAAACAACAGGACAGTGCCAGAACTTTCACGTTGAAAACAACAGGACAGTGTTAGTTTTTTTT	3930	TGTCCATCAT		4025	CAGTCACTGAT	4100	rggaatctctg	4175	<i>PEAGTGGCGG</i>		4250	CAGTGGACCTC	4325	AAGTCCATGAG	4400	SCAGTTCCCCA		4475	AACCTCTATGC	4550	SAACAGGGAGT	
GTGGAGCAGGGAAGGGTATGGATGTAAA BSPMI- 3985 3995 4005 4060 4070 4080 TTGCTGGAAACCCAGAGTTTCACGTTGAAAAC 4135 AGATCTGCTAACAGCTTGGTTGAAAAC Bglil 4210 4220 4230 CCCCGCTATTCCTTACTTGCTGTAGTTGTTA 4285 4295 4305 GGTCCAAGTCCCTGTCCTATTTTACTCTTGTA 4285 44295 4436 4445 4445 4445 AATGTCGGATCTGGCCCCTTTACTTGTTACTTACTTACTT	3940	GAGTTTGAAAT		4015	AACAGAGTGC	4090	AACAGGACAG 1	4165	TCAGCTTAGG		4240	TTTTGCTTAT	4315	AGGTGCAGAGA	4390	AGTGCACAGG	ApaLI	4465	ATAATGTGAA	4540	AAAGTGCTTA	
GTGGAGCAGGTGGGAAGGGTA1 BSPMI- 3985 GGCCTATGTCAGCAGTCACAGCC 4060 4070 TTGCTGGAACCCAGAAGTTTCA 4135 AGATCTGCTAACACTGATCTTGC Bg11I 4210 4285 GAGTCTATGTCCCTATTGTCTTA 4360 GGTCCAAGTCAGGGACACTATTC 4435 AATGTCGGATCTGGCCCCTTCCT 4510 ACAGGGATAATCCCAGAACTGA(3930	GGATGTGTAA		4005	TGGAGGTGGT	4080	ACGTTGAAAAC	4155	STIGGCIGCCG		4230	CCTCGGTCTA	4305	CATCCTCTGTA	4380	CAGCAATCTAC		4455	FTCCCCACTGT	4530	STTGTCCATGT	
GTGGAGCAGGTG BSPMI- 3985 3985 GGCCTATGTCAG 4060 TTGCTGGAAACC 4135 AGATCTGCTAAC Bg11I 4210 CCCCGCTATTCC 4285 GAGTCTATGTGT 4360 GGTCCAAGTCAG 4350 AATGTCGGATCT AA35 AATGTCGGATCT ACAGGGATAATC	3920	GGGAAGGGTAI			CAGTCACAGCC		CAGAAGTTTCA		ACTGATCTTG			CTATTGTCTT		CCCTGTCCCAC		GGACACTATT			GCCCCTTCC		CCAGAACTGAG	
	3910	GTGGAGCAGGTG	BspMI-	3985	GGCCTATGTCAG	4060	TTGCTGGAAACC	4135	AGATCTGCTAAC	BglII	4210	CCCCCCTATTCC	4285	GAGTCTATGTGT	4360	GGTCCAAGTCAG		4435	AATGTCGGATCT	4510	ACAGGGATAATC	

FIG. 1A-6

GAAACATGGTGGTCCGGGCCTGTGGCTGCCACTAGCTCCTCCGA

FIG. 1B

CONSENSUS PROBE 20 30 40 50 60 70 GATCCTAATGGGCTGTACGTGGACTTCCAGCGCGACGTGGGACGACTGGATCATCGCCCCCGTCG ** *** *** *** *** *** ****** ** TGTAAGAAGCACGAGCTTCCGAGACCTGGGCTGGCAGGACTGGATCATCGCGCCTGAAG OP4 28 88	ACTTCGACGCCTACTACTGCTCCGGAGCCTGCCAGTTCCCCTCTGCGGATCACTTCAACAGCACCAACCA	150 160 210 210 210 210 210 210 210 210 210 21	220 230 240 250 260 270 280 GAGCTGTCCGCCATCAGCATGCTGACCTGACCAGGAGA **** **** *** *** *** *** *** CAGCTCAATGCCATCCTCATCCAGGAGAATTCCAACGTGCTGAAGAACTACCAGGAGA **** *** *** *** *** *** *** *** *** *	310 CTGCCGCTAACTGCA **** ** ** CTGCCACTAGCTCCT 328
20 30 CGTGGACTTCCAGCGC ** CGTATGTCAGCTTCCGA 38 48	90 100 TECTCCGGAGCCTGCC ** ** TGTGAGGGGGAGTGTG	160 CCTGGTGAACAACATGAA **** ** ** ** CTGGTCCACTTCATCAA	230 240 SCATGCTGTACCTGGAC ** ** ** CGTCCTCTACTTCGAT 248 258	300 310 GCGGCTGCCGCTAACTGC * ***** ** GTGGCTGCCACTAGCTCC 318 328
CONSENSUS PROBE 20 GATCCTAATGGGCTGTACGT ** TGTAAGAAGCACGAGCTGTA OP4 28 38	80 90 FTCGACGCCTACTACTGC * ** ****** FACGCGCCTACTACTG1	150 CGCCGTGGTGCAGACCCTGG **** **** **** CGCCATCGTGCAGACGCTGG	220 AGCTGTCCGCCATCAGCAT **** ****** AGCTCAATGCCATCTCCGT 238 248	290 310 TGACCGTGGGCTGCGGCTGCCGCTAACTGCA ** ** ** ** ** ** TGGTGGTCCGGCCTGTGGCTGCCACTAGCTCCT 308 318



------ CUFFT

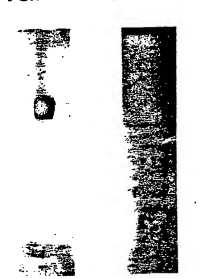


FIG. 5A





FIG. 3A FIG. 3B FIG. 4A FIG. 4B

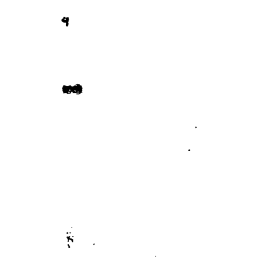


FIG. 5B



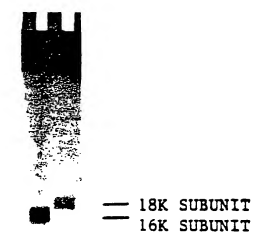
WO 89/09787

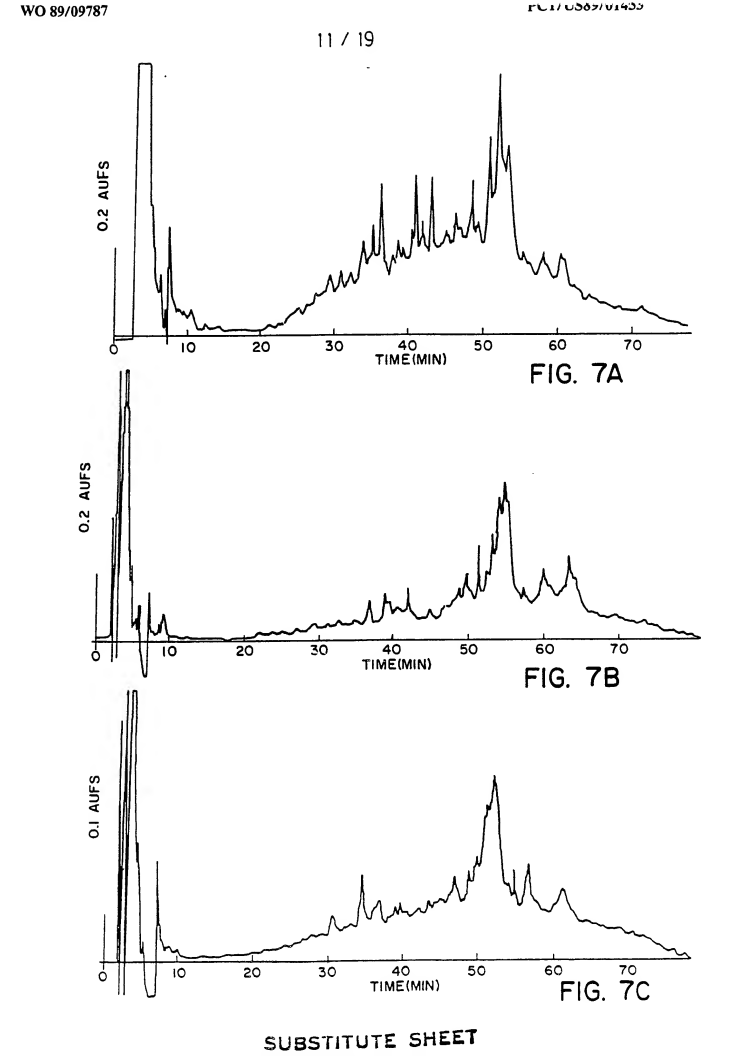
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TC1/U569/U1453

FIG.6 A FIG.6 B FIG.6 C FIG.6 D FIG.6 E







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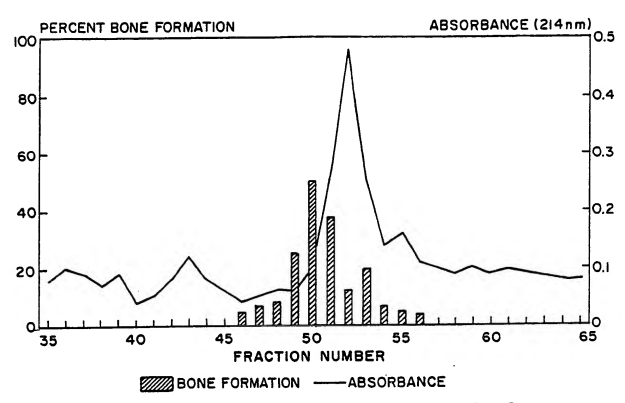


FIG. 8

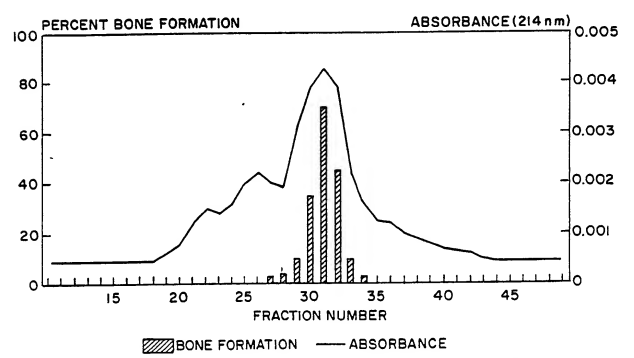
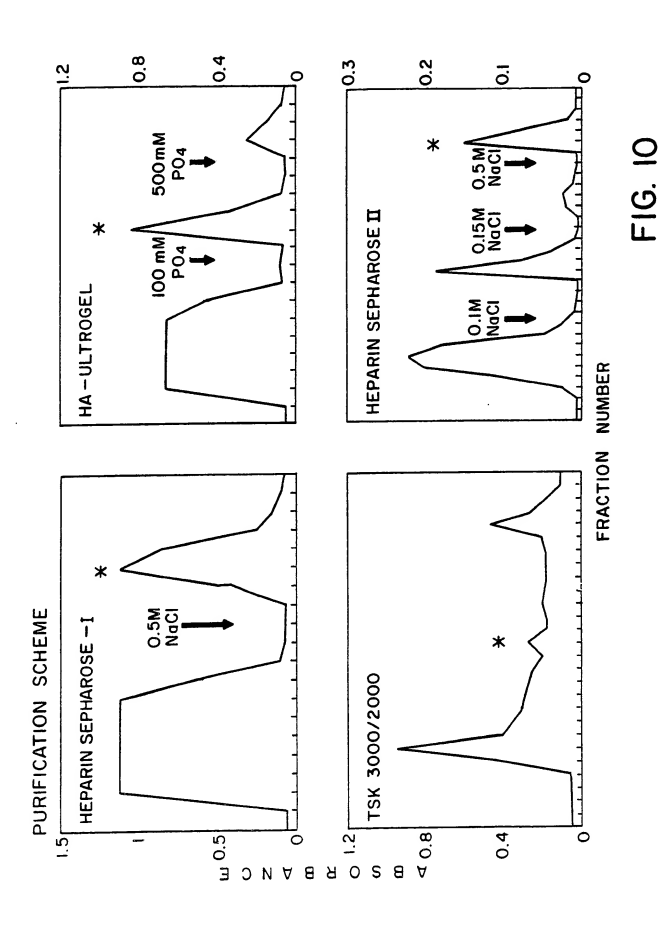
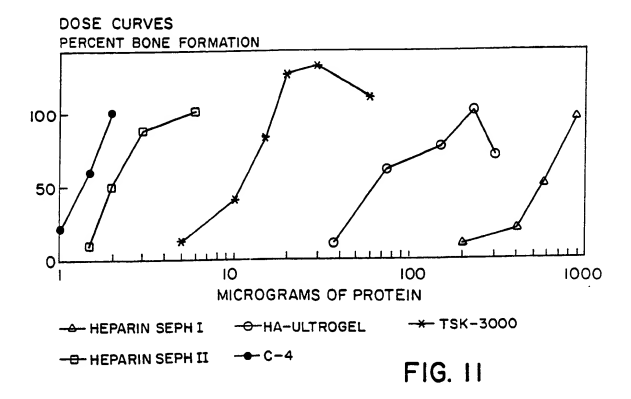


FIG. 9

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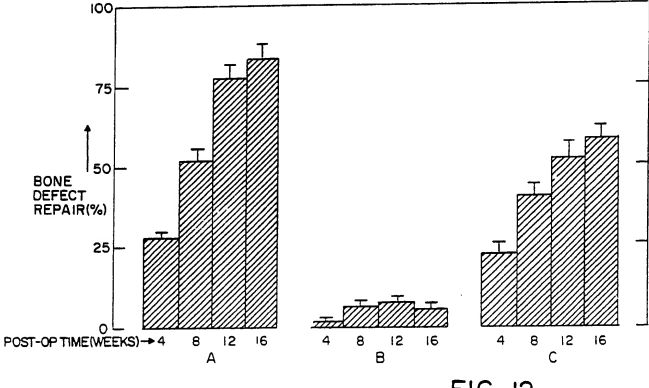


FIG. 12

FIG. 13

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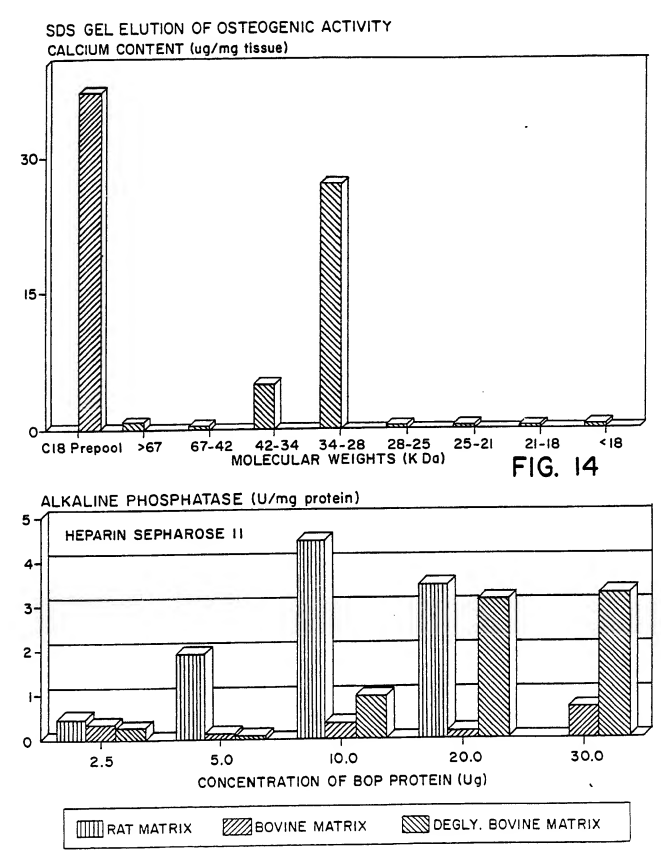
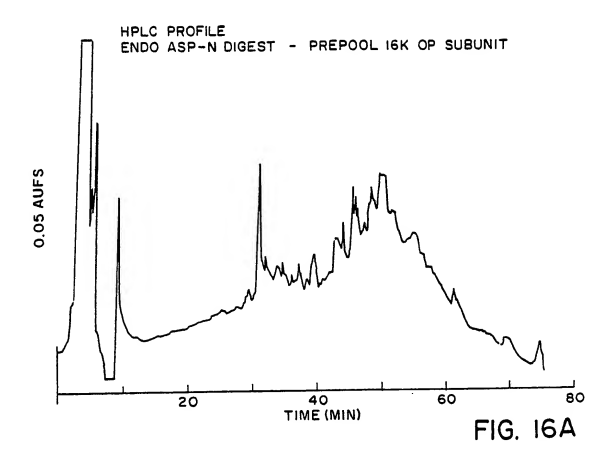


FIG. 18

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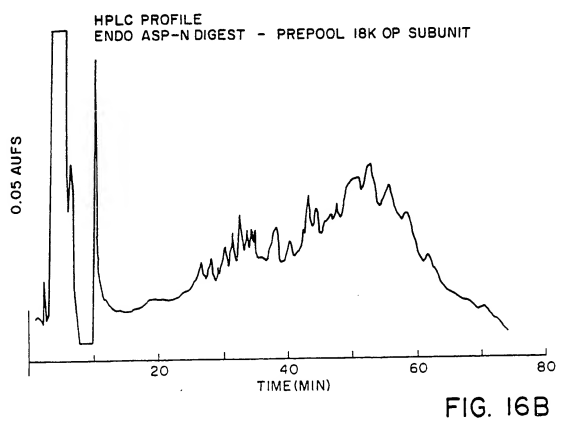




FIG. 17A

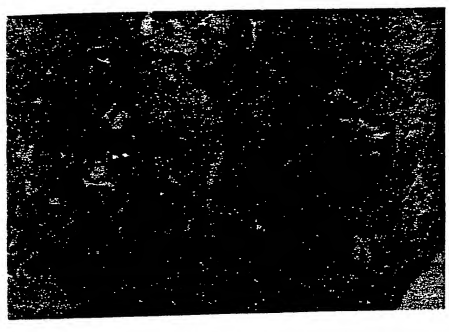


FIG. 17 B

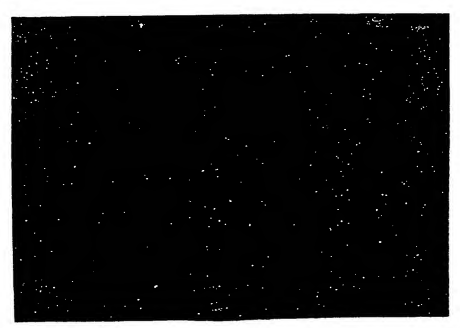


FIG. 17C

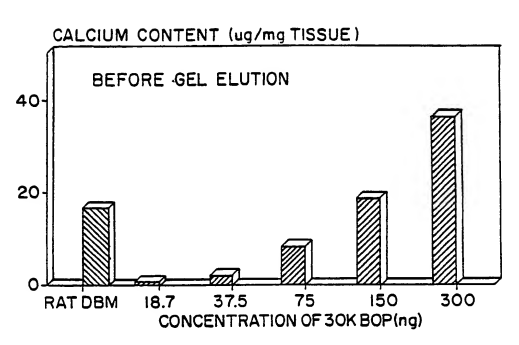


FIG. 19A

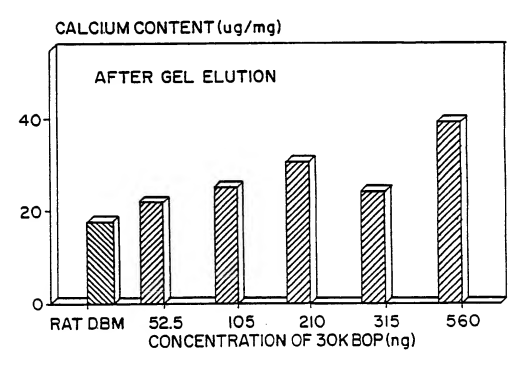


FIG. 19B

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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

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C07K 13/00, A61L 27/00, C12N 15/00
A61K 35/32, C12P 21/02

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 US

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 15 August 1988 (15.08.88)
 US

 315,342
 23 February 1989 (23.02.89)
 US

(60) Parent Applications or Grants

(63) Related by Continuation
US
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(71) Applicant (for all designated States except US): CREATIVE BIOMOLECULES, INC. [US/US]; 35 South Street, Hopkinton, MA 01748 (US).

(72) Inventors; and

(75) Inventors, and (75) Inventors, Applicants (for US only): KUBERASAMPATH, Thangavel [IN/US]; 6 Spring Street, Medway, MA 02053 (US). OPPERMANN, Hermann [US/US]; 25 Summer Hill Road, Medway, MA 02053 (US). RUE-GER, David, C. [US/US];

150 Edgemere Road, Apt. 4, West Roxbury, MA 02132 (US). OZKAYNAK, Engin [TR/US]; 44 Purdue Drive, Milford, MA 01757 (US).

(74) Agent: PITCHER, Edmund, R.; Lahive & Cockfield, 60 State Street, Boston, MA 02109 (US).

(81) Designated States: AT (European patent), AU, BB, BE (European patent), BF (OAPI patent), BG, BJ (OAPI patent), BR, CF (OAPI patent), CG (OAPI patent), CH (European patent), CM (OAPI patent), DE (European patent), DK, FI, FR (European patent), GA (OAPI patent), GB (European patent), HU, IT (European patent), JP, KP, KR, LK, LU (European patent), MC, MG, ML (OAPI patent), MR (OAPI patent), MW, NL (European patent), NO, RO, SD, SE (European patent), SN (OAPI patent), SU, TD (OAPI patent), TG (OAPI patent), US.

Published

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(88) Date of publication of the international search report: 8 February 1990 (08.02.90)

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INTERNATIONAL SEARCH REPORT

International Application No.

PCT/US 89/01453

1. CLASSIFICATION OF SUBJECT MATTER (if several classification symbols apply, indicate all) 4				
According to International Patent Classification (IPC) or to both National Classification and IPC				
IPC4: C 07 K 13/00, A 61 L 27/00, C 12 N 15/00, A 61 K 35/32,				
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Documentation Searched other than Minimum Documentation to the Extent that such Documents are included in the Fields Searched *				
III. DOCUMENTS CONSIDERED TO BE RELEVANT				
Category •	Citation	of Document, 11 with Indication, wher	appropriate, of the relevant passages 12	Relevant to Claim No. 13
х,у	wo,	A, 88/00205 (GEN) 14 January 1988 see pages 1-12,19 pages 61-73, class	5-17,22-24.49:	1-16,25, 26,28,32, 33
Y	The	The Journal of Cell Biolgoy, volume 97 December 1983, The Rockefeller University Press, S.M. Seyedin et al.: "In vitro induction of cartilage-specific macromolecules by a bone extract", pages 1950-1953 see the whole article, especially page 1952, right-hand column		
Y	EP,	A, 0182483 (COLLAGEN CORP.) 28 May 1986 see the whole document, especially page 6, first paragraph; page 7, lines 10-18		1-16,25, 26,28,32, 33
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Date of the Actual Completion of the Leavest				
Date of the Actual Completion of the International Search 13th November 1989 4.0.04.00				
International Searching Authority Signature of Authorized Offices				
EUROPEAN PATENT OFFICE T.K. WILLIS				

Category • .	Citation of Document, with indication, where appropriate, of the relevant passages	r Relevant to Claim No
Y	WO, A, 85/05274 (R.F. OLIVERS) 5 December 1985 see the whole document	! 8
Y :	Analytical Biochemistry, volume 146, 1985, Academic press, Inc., C.A. Olson et al.: "Deglycosylation of chondroitin sulfate protecglycan by hydrogen fluoride in pyridine", pages 232-237 see "Discussion" on page 236	8
A	Trends in Biochem. Sci. (TIBS), volume 9, 1984, Elsevier Science Publishers B.V., (Amsterdam, NL),. E. Simpson: "Growth factors which affect bone", pages 527-530 see the whole article	1-19,23-33
A	EP, A, 0148155 (DOW CHEMICAL CO.) 10 July 1985 cited in the application	
A	WO, A, 86/00526 (A.I. CAPLAN) 30 January 1986	
A :	US, A, 4394370 (S.R. JEFFERIES) 19 July 1983	
A	US, A, 4563489 (M.R. URIST) 7 January 1986	
A :	S.P. Colowick et al.: "Methods in Enzymology", volume 146, Peptide Growth Factors, part A, edited by David Barnes et al., Academic Press Inc., M.R. Urist et al.: "Preparation and bioassay of bone morphogenetic protein and polypeptide fragments", pages 294-312	
Α	EP, A, 0169016 (COLLAGEN CORP.) 22 January 1986	
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FURTHER INFORMAT	ION CONTINUED FROM THE SECOND SHEET	
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V. OBSERVATIONS	WHERE CERTAIN CLAIMS WERE FOUND UNSEARCHABLE	
This international search of		
1. Claim numbers **	eport has not been established in respect of certain claims under Article 17(2) (a) for the following read * because they relate to subject matter not required to be searched by this Authority, namely:	sons:
See Por Pu	numbers 20-22: ule 39.1(iv)	
Methods for	or treatment of the human or animal body	
by surgery	y or therapy, as well as diagnostic methods.	
	and the designation of the children in the chi	
2. Claim numbers ** ments to such an exte	K., because they relate to parts of the international application that do not comply with the prescribed tent that no meaningful international search can be carried out, specifically:	require-
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See Articl	le 6 PCT, 17(2)(a)(ii) PCT	
	./.	
Claim numbers	Page up the control of the control o	
PCT Rule 6.4(a).	, because they are dependent claims and are not drafted in accordance with the second and third sente	nces of
VI. X OBSERVATIONS	WHERE UNITY OF INVENTION IS LACKING 2	
	g Authority found multiple inventions in this international application as follows:	
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	101. 101/15A/208 dated 14-8-89	
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As all required addition	nal search fees were timely paid by the applicant, this international search report covers all searchable plication.	claims
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those claims of the inte	equired additional search fees were timely paid by the applicant, this international search report cover ernational application for which fees were paid, specifically claims:	rs only
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the invention first ment	search fees were timely paid by the applicant. Consequently, this international search report is restrictioned in the claims; it is covered by claim numbers:	cted to
As all searchable claims	s could be searched without effort justifying an additional fee, the International Searching Authority of	a:a = - •
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_	ees were accompanied by applicant's protest.	
No protest accompanie	d the payment of additional search fees.	

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The present application is admittedly <u>NOT</u> the first to describe the phenomenon of bone inducing proteins, or devices containing them. It is therefore necessarily drafted to the <u>individual compounds</u> (on well-known devices).

In claims 1-16, however, proteins (on a device) are claimed, which are only defined by their biological activity and/or their molecular weight, which is clearly not sufficient for a full characterization of individual compounds. Despite this broad scope, the biological activity has only been demonstrated for one single compound (BOP 30K, which is covered by subject 1).

The search has therefore been restricted to the embodiments of claims 1-19 and 23-33, in as far as the proteins (and DNAs) correspond to the definitions given in the claims 17, 18, and 23-31.

This searchable subject matter has been regrouped according to the non-unity specification, in order to establish conceptually individual subjects, each of which now constitutes a potential selection invention.

ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

US 8901453 SA 28156

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 19/12/89

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(71) Applicant (for all designated States except US): CREATIVE BIOMOLECULES, INC. [US/US]; 35 South Street, Hopkinton, MA 01748 (US).

(72) Inventors; and

(75) Inventors, and
(75) Inventors/Applicants (for US only): KUBERASAMPATH,
Thangavel [IN/US]; 6 Spring Street, Medway, MA
02053 (US). OPPERMANN, Hermann [US/US]; 25
Summer Hill Road, Medway, MA 02053 (US). RUEGER, David, C. [US/US];

150 Edgemere Road, Apt. 4, West Roxbury, MA 02132 (US). OZKAYNAK, Engin [TR/US]; 44 Purdue Drive, Milford, MA 01757 (US).

- (74) Agent: PITCHER, Edmund, R.; Lahive & Cockfield, 60 State Street, Boston, MA 02109 (US).
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INTERNATIONAL SEARCH REPORT

International Application No

PCT/US 89/01453

I. CLAS	SIFICATION	OF SUB	JECT MA	TTER	(it se	veral cla	ssifics	tion s	ymbo	is ac	niv.	Indic	ate a	ID 6				,, 0.	T 4 7
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		Minimum Documentation Searched 7 Classification Symbols C 07 K, C 12 N, A 61 K, A 61 L, C 12 P Documentation Searched other than Minimum Documentation to the Extent that such Documents are included in the Fields Searched 8 FS CONSIDERED TO BE RELEVANT* Citation of Document, 11 with indication, where appropriate, of the relevant passages 12 WO, A, 88/00205 (GENETICS INSTITUTE) 14 January 1988 See pages 1-12,15-17,22-24,49; pages 61-73, claims 1-23 The Journal of Cell Biolgoy, volume 97. December 1983, The Rockefeller University Press, S.M. Seyedin et al.: "In vitro induction of cartilage-specific macromolecules by a bone extract", pages 1950-1953 See the whole article, especially page 1952, right-hand column EP, A, 0182483 (COLLAGEN CORP.) 28 May 1986 See the Whole document, especially page 6, first paragraph; page 7, lines 10-18 This set cited documents: 16 Principle of the pri																	
Category •	Citation	of Docum	nent, 11 wi	th Indic	ation,	where a	ppropr	iate, o	f the	relev	ent p	2888	ges '	2	Re	levan	t to C	aim No	0. 13
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Category * .	Citation of Document, with indication, where appropriate, of the relevant passages	. Relevant to Claim No
Y :	WO, A, 85/05274 (R.F. OLIVERS) 5 December 1985 see the whole document	! 8
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A :	EP, A, 0169016 (COLLAGEN CORP.) 22 January 1986	

FURTHER INFORMATION CONTINUED FROM THE SECOND SHEET	37,02133
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V. OBSERVATIONS WHERE CERTAIN CLAIMS WERE FOUND UNSEARCHABLE	
This international search report has not been extended in	
This international search report has not been established in respect of certain claims under Article 17(2) ** Claim numbers	(a) for the following reasons: Authority, namely:
** Claim numbers 20-22:	1
See PCT Rule 39.1(iv)	•
Methods for trootmont of the	
Methods for treatment of the human or animal	body
by surgery or therapy, as well as diagnostic	methods.
• • • • • • • • • • • • • • • • • • • •	
2. Claim numbers <u>**</u> , because they relate to parts of the international application that do not comments to such an extent that no meaningful international search can be carried out, specifically:	ply with the prescribed require-
** Claim numbers 1-16,19,32,33	
See Article 6 PCT, 17(2)(a)(ii) PCT	
C 1c1, 1/(2)(a)(11) PCT	
./.	
Claim numbers, because they are dependent claims and are not drafted in accordance with the PCT Rule 6.4(a).	
PCT Rule 6.4(a).	SECOND END THIS SENTENCES OF
VI. X OBSERVATIONS WHERE UNITY OF INVENTION IS LACKING 2	
This international Searching Authority found multiple inventions in this international application as follows	:
Please see Form PCT/ISA/206 dated 14-8-89	·
= +4, -5 , 250 dacta 14 -6-69	
As all required additional accept to a superior of the superio	
As all required additional search fees were timely paid by the applicant, this international search report of the international application.	t covers all searchable claims
As only some of the required additional search fees were timely gold but to gold but to	
those claims of the international application for which fees were paid, specifically claims:	nai search report covers only
• "	
No required additional accept to	
	search report is restricted to
the invention first mentioned in the claims; it is covered by claim numbers:	
As all searchable claims could be searched without office lives	
As all searchable claims could be searched without effort justifying an additional fee, the International invite payment of any additional fee.	Searching Authority did not
emark on Protest	
The additional search fees were accompanied by applicant's protest.	
No protest accompanied the payment of additional search fees.	j

FURTHER INFORMATION CONTINUED FROM PCT/ISA/210 (supplemental sheet (2))

The present application is admittedly <u>NOT</u> the first to describe the phenomenon of bone inducing proteins, or devices containing them. It is therefore necessarily drafted to the <u>individual</u> <u>compounds</u> (on well-known devices).

In claims 1-16, however, proteins (on a device) are claimed, which are only defined by their biological activity and/or their molecular weight, which is clearly not sufficient for a full characterization of individual compounds. Despite this broad scope, the biological activity has only been demonstrated for one single compound (BOP 30K, which is covered by subject 1).

The search has therefore been restricted to the embodiments of claims 1-19 and 23-33, in as far as the proteins (and DNAs) correspond to the definitions given in the claims 17, 18, and 23-31.

This searchable subject matter has been regrouped according to the non-unity specification, in order to establish conceptually individual subjects, each of which now constitutes a potential selection invention.

ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

US 8901453 SA 28156

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 19/12/89

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